BLUMENTHAL FOODY WONG

PREVENTIVE CARDIOLOGY

A Companion to Braunwald's

HEART DISEASE



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A Companion to Braunwald's Heart Disease



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This book is dedicated to the memory of Dr. Kenneth L. Baughman, who exemplified a tremendous commitment and personal passion for the principles and teachings of preventive cardiology during his entire life.

We would also like to thank our families for their support and encouragement during the development of this comprehensive textbook.

In addition, we extend special appreciation to those who inspired our careers in preventive cardiology, namely Drs. Eugene Braunwald, Peter Libby, Thomas Pearson, Adrian Ostfeld, William Kannel, William Castelli, Jeremiah Stamler, and Peter Kwiterovich.

Finally, we remember key colleagues and friends, including Dr. Stanley Blumenthal, Henry Ciccarone, David Kurtz, and John Yasuda, who made a difference in our lives and our commitment to preventive cardiology.

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Computed Tomography in Evaluation and Prevention of
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Foreword

persons were struck down during their most productive years, and be used in refining the assessment of coronary risk. at a time of large family responsibilities. These "heart attacks" were rarely considered.

called the "father of American cardiology." As early as the 1930s, broad audience. Prevention of cardiovascular disease is too White always included a section on prevention in his lectures on important to leave to a relatively small group of experts, but instead coronary artery disease, and he wrote about it in his famous must be carried out by all physicians, regardless of specialty, as well textbook. The National Heart Institute (now the Heart, Lung and as by nurses and other health care professionals who care for Blood Institute) was established in 1948 and was instrumental in patients with, or at risk of developing, cardiovascular disease. All of furthering the concept of cardiac disease prevention. Two of the these groups and their trainees can profit enormously from this most important early actions by the Institute were the establishment important book. of the Framingham Heart Study and of the Lipid Research Clinics. The former was (and continues to be) a long-term prospective study, to the growing list of Companions to Heart Disease. with standardized examinations at intervals of adults who were initially without clinical manifestations of coronary artery disease. By 1961, it was evident that overtly healthy subjects with hypertension, hypercholesterolemia, and/or who were cigarette smokers were at higher risk to develop acute myocardial infarction than were their age- and sex-matched controls without these characteristics. Framingham investigators thus coined the term Peter Libby "coronary risk factors." These observations led to the important idea that the amelioration of risk factors would prevent, or at least delay, the development of clinical coronary artery disease. Considerable research has been done during the past half century that has supported this idea.

The institute's second major contribution was the Coronary -Primary Prevention trial, which demonstrated that in subjects with hypercholesterolemia, but without overt coronary artery disease, the occurrence of coronary events could be reduced with a diet and cholestyramine, a resin that reduces elevated serum cholesterol. This confirmed, once and for all, the important role of cholesterol in atherogenesis. A breakthrough in coronary prevention occurred in the 1980s with the development of HMGCoA reductase inhibitors (statins), which caused a substantial lowering of LDL-cholesterol. Simultaneously, well tolerated blood pressure-reducing drugs and smoking cessation programs were developed.

At first, many cardiologists reacted sluggishly to these observations and often did not incorporate preventive measures into their practices. Both the glamor (and reim bursement) favored the diagnosis and management of acute illness over the more mundane (and poorly reimbursed) efforts required to maintain patientsparticularly those who had no overt cardiovascular disease - on diet and other life style measures as well as drugs, which often have some annoying side effects. However, during the 1990s, the evidence in favor of the clinical benefits of prevention became overwhelming, and in the first decade of this century, expert committees developed practical guidelines that provided strong support. Adherence to these guidelines became important measures of physician performance, a trend that only promises to increase in coming years.

Now, in the second decade of the current century, preventive cardiology has a robust and rapidly growing knowledge base. In addition to hypercholesterolemia, hypertension and cigarette

In the middle of the twentieth century, the development of an acute smoking described half a century ago, we now recognize that myocardial infarction was often totally unexpected and like the diabetes, vascular inflammation, kidney disease, passive smoking, proverbial "bolt out of the blue." Frequently, apparently healthy and a growing number of biomarkers and genetic variants may also

Preventive Cardiology is very capably edited by Drs. Blumenthal, often either fatal or disabling. Medical attention was focused largely Foody, and Wong, and written by stellar authors, all experts in their on the diagnosis and management of these catastrophic events. subjects. It is a superb, well written and illustrated volume that Forestall ing or even better, preventing, myocardial infarction was elegantly weaves together the many separate strands of this critically important area of cardiology to provide a thorough One notable exception, however, was Dr. Paul D. White, often understanding of the field. This volume should serve the needs of a

We are therefore very pleased to welcome *Preventive Car diology*

E ugene Braunwald ROBERT BONOW Douglas Mann DOUGLAS ZIPES

For nearly a century, atherosclerotic cardiovascular disease has remains clinically silent for decades before resulting in an acute ischemic syndrome, myocardial infarction, stroke, or sudden cardiac death. Since atherosclerosis is a progressive disease that starts early in life, it challenges us to be more aggressive in our efforts regarding prevention.

improve people's quality of life. Unfortunately, rates of obesity and related conditions such as metabolic syndrome and diabetes are on and e motional aspects of preventive cardiology. the rise, in both developed and developing countries. Instead of prevention, significant health care dollars are spent on the end-stage prevention of cardiovascular disease. It provides an overview of the complications of atherosclerotic vascular disease, such as drug- epidemiology and risk factors for cardiovascular disease, and the eluting stents, implantable cardioverter-defibrillators, and surgical revascularization.

emphasize preventive strategies to slow or halt the progression of tools provided in this text we may achieve the promise of the atherosclerosis. Health care providers need to understand how to prevention of most cardiovascular disease events in our lifetimes. optimize cardiovascular risk stratification. The Framingham and other global risk algorithms serve as an important starting point in risk assessment, but have limitations and often exclude key risk factors such as a family history of premature cardiovascular disease, glucose intolerance, triglycerides, waist size, and lifestyle habits. For example, although an adult with a glucose level of 126 mg/dL or higher is automatically placed into a very high risk category, a similar individual with a slightly lower glucose level but who may have additional risk factors or evidence of advanced subclinical atherosclerosis for their age may actually be at higher risk, but would not necessarily qualify for aspirin therapy, antihypertensive therapy, or lipid-lowering therapy.

A great need also exists for better understanding of the significance, clinical utility, and cost-effectiveness of more novel risk factors and screening for asymptomatic cardiovascular disease. Atherosclerosis imaging and measurement of biomarkers such as hs-CRP are now fairly widely performed, and there is a need for understanding how to incorporate into clinical practice the findings from large-scale epidemiological studies (eg, Cardiovascular Health Study and the Multi Ethnic Study of Atherosclerosis) and clinical trials such as JUPITER. However, there are clear limitations to the data that we have so far on biomarkers such as hs-CRP and increasingly popular multimarker approaches, and imaging measures such as coronary artery calcium and carotid intima-media thickness. Experts are clearly split on how to incorporate emerging risk factors and subclinical disease into clinical practice.

The medical community needs to promote guideline adherence and reduce the gap in the use of proven medical and lifestyle therapies. Moreover, federal, state, and local governments, education departments and schools, and the corporate sector need to play a greater role in ensuring environments conducive to promoting heart health. The cornerstone of prevention is based on therapeutic lifestyle changes, including regular brisk physical activity and a healthy diet, and strategies to better support these measures need to be developed and implemented at the health care and community level.

In this companion to Braunwald's Heart Disease, we approach cardiovascular disease prevention in a convenient ABCDE framework. In 2002 the AHA and ACC produced a guideline statement on the management of patients with chronic stable angina and arranged their recommendations into an ABCDE format. This approach has also been used as the basis for the training of fellows in preventive cardiology. It has also been used in several evidencebased reviews on primary and secondary prevention of CVD, management of non-ST-segment elevation myocardial infarction (NSTEMI) and management of metabolic syndrome. 2-4

Prevention needs to be a central feature of a sustainable health

care system, but implementation of preventive practices remains been the leading cause of death in industrialized countries. It often suboptimal. The ABCDE approach arranges prevention guidelines into an easy-to-remember framework that can be used by clinicians with each patient to ensure comprehensive care. The main sections of this textbook include: (A) an assessment of risk from a clinical and genetic perspective, a therothrombosis and antiplatelet therapy; (B) **blood** pressure management; (C) **cholesterol** and dyslipidemia; (D) Early identification of cardiovascular risk and modification of diet and lifestyle issues (diabetes mellitus, metabolic syndrome; risk factors reduce the incidence of future cardiovascular events and disparities in care; diagnostic testing to help improve risk prediction); and (E) e xercise prescriptions, cardiac rehabilitation

This text is meant to serve as a guide for those interested in the importance of risk stratification. It under scores the evidence base for the management of cardiovascular risk factors and provides Physicians, nurses, and other health care providers need to recommendations for clinical care. It is our hope that armed with the

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SECTION |

Risk Assessment

CHAPTER 1

Preventive Cardiology: Past, Present, and Future

KEY POINTS

Michael J. Blaha, Ty J. Gluckman, and Roger S. Blumenthal

- Atherosclerotic cardiovascular disease (CVD) is an ideal scenario for prevention efforts because (1) it is a common disease; (2) it is modifiable by behavior; (3) it has a long latency; (4) the time between symptom onset and severe disability or sudden cardiac death is short; and (5) no cure exists for systemic atherosclerosis once it is present.
- The Framingham Heart Study identified smoking, elevated blood pressure, and high cholesterol as the principal risk factors for CVD. More recently, the INTERHEART study has shown that 9 main CVD risk factors account for 90% of the population-attributable risk for a first myocardial infarction.
- The majority of improvement in rates of mortality from CVD since the 1960s is the result of prevention, not treatment, of acute CVD.
- Prevention occurs at three levels: primordial, primary, and secondary. However, there may be variable degrees of overlap as the cutoff points for risk factors change and as imaging modalities identify populations with disease burden that is not expected on the basis of traditional risk factors.

- There are two main approaches to prevention: a population-based approach, in which researchers seek to make small changes in risk factors across the entire population, and an individual-based approach, which emphasizes identifying individuals at high risk for CVD and aggressively lessening their risk factors.
- Guideline and scientific statements from the American Heart Association (AHA), American College of Cardiology (ACC), and others

Since the early 1900s, atherosclerotic - cardiovascular disease (CVD), including both coronary heart disease (CHD) and stroke, has been the leading cause of death in industrialized nations. ¹ Atherosclerosis represents a unique public health challenge because it is a progressive, lifelong disease that is modified by behavior and yet - produces few symptoms until late into its course. Unfortunately, when it does become clinically evident, there is often a short duration between symptom onset and - disability, and sudden death is a common sen tinel event.

In spite of numerous advances that have improved the treatment of acute CVD, many therapies remain costly, and their effectiveness depends on the prompt identification of the few individuals most likely to benefit. Both reperfusion and revascularization procedures are indicated in only a select group of patients with critical occlusive vascular disease; these treatments target localized areas of the vascular bed without

direct population-based and individual-based preventive care organizations.

- Despite guidelines, there is a wide gap between the burden of CVD and current preventive efforts.
 This gap can be narrowed with more simplified, comprehensive guidelines.
- This chapter offers an easy-to-remember memory tool that facilitates comprehensive preventive care: the "ABCDE" approach.

addressing atherosclerosis throughout the rest of the body. As such, there remains no cure for atherosclerosis as a systemic disease.

Nevertheless, disproportionately large amounts of money are spent late in the disease course on relatively small numbers of patients with acute complications of CVD, rather than the far greater numbers in whom early preventive efforts might lead to markedly greater benefit. These factors underscore the true importance of CVD prediction and prevention, and they preface not only this chapter but the content of this entire text on preventive cardiology (Box 1-1).

With great foresight, the US Public Health Service launched a publicly funded effort in the 1940s to identify modifiable CVD risk factors. Through modern clinical epidemiologic methods, the landmark Framingham Heart Study ²helped define the field of preventive cardiology and led to the identification of smoking, hypertension,

BOX 1-1 Factors Making Atherosclerosis Ideal for Prevention

- High incidence
- Modifiable by behavior
- Long disease latency
- Short time between symptoms and disability
- Sudden death: a common manifestation
- Available treatments unable to cure the underlying disease
- Treatment of acute diseases associated with huge financial and societal costs

and elevated cholesterol as the "main risk factors" for CVD. ³ In the years that followed, the US government launched several population-based educational campaigns and spent billions of dollars funding research aimed at controlling these risk factors. The Atherosclerosis Risk in Communities (ARIC) study, the Coronary Artery Risk Development in Young Adults (CARDIA) study, the Cardiovascular Health Study (CHS), and the Multi-Ethnic Study of Atherosclerosis (MESA) were instrumental in the effort to identify novel risk factors, to describe the determinants of early Apo, apolipoprotein; CI, confidence interval. atherosclerosis, and to understand these factors and determinants in relation to younger, older, and multiple ethnic populations. Unfortunately, in spite of these efforts, smoking, hypertension, and hypercholesterolemia remain unacceptably common in the general population today. 1

Risk factors for CVD begin accumulating at a young age, often while individuals are asymptomatic and unaware of the untoward consequences. Pathological evidence of atherosclerosis can be identified soon after risk factor onset; persons with measurable risk demonstrate this evidence earliest. 4,5 Although risk factors are frequently present as early as the second and third decades of life, the presence of multiple risk factors is associated with an even higher prevalence of early atherosclerotic vascular disease. 6 Never has the risk for such individuals been more important than it is today, when a burgeoning global epidemic of childhood obesity further heightens the public health challenge.

Results from the global INTERHEART study suggest that nine modifiable risk factors - dyslipidemia, smoking, diabetes mellitus, hypertension, abdominal obesity, psychosocial stress, poor diet, physical inactivity, and alcohol consumption -account for more than 90% of the risk for a first myocardial infarction (Table 1-1). 7 The effects of these risk factors appear to be remarkably stable across gender, race, and geographic location. Such data have led the World Health Organization (WHO) to estimate that 80% of premature CHD can be prevented with comprehensive assessment and management of these risk factors. 8

Because major CVD risk factors often co-occur, emerging risk factors probably account for disproportionately smaller numbers of CVD events. ⁹ In epidemiological terms, biomarkers such as interleukin-6, adiponectin, and lipoprotein(a) are associated with a campaigns to decrease salt intake have resulted in significant smaller incremental population-attributable risk. The value of reductions in systolic blood pressure. 19,20 measuring these factors, therefore, lies more in elucidating the pathophysiological mechanisms of CVD and identifying novel therapeutic targets than in global risk prediction (Box 1-2).

Much research is still needed to better integrate existing risk variables into prediction models of short- and long-term global risk. This is important not only to ensure the cost effective use of existing risk-reducing therapies (eg, aspirin and statins) but also to determine who may benefit from measurement of biomarkers or 47% resulted from evidence-based medical therapy directed at detection of subclinical atherosclerosis through imaging patients with known or suspected vascular disease. 21 Importantly, techniques. ¹⁰ Improved treatment decisions – including delivery of existing options and the selective use of new modalities – remains the mainstay of

TABLE 1-1	ABLE 1-1 Interheart: A Global Case-Control Study of Risk Factors for Acute Myocardial Infarction			
Risk Factor	Odds Ratio (99% CI) Multivariable Adjusted	Population-Attributable Risk Multivariable Adjusted		
ApoB/ApoA-I	3.25 (2.82-3.76)	49%		
Current smoking	2.87 (2.58-3.19)	36%		
Diabetes	2.37 (2.07-2.71)	9.9%		
Hypertension	1.91 (1.74-2.10)	18%		
Abdominal obesity	1.62 (1.45-1.80) 2.67 (2.21-3.22)	20% 33%		
Psychosocial stress and depression				
Daily fruit and vegetable intake	0.70 (0.62-0.79)	14%		
Exercise	0.86 (0.76-0.97)	12%		
Alcohol intake	0.91 (0.82-1.02)	7%		
Combined	129	90%		

BOX 1-2 Modern Themes in Cardiovascular Disease Risk Prediction

Novel risk factors: increasingly diminished population-attributable risk

Novel risk factors: value is likely to be weighed in elucidating pathophysiological mechanisms and guiding treatment

Need for improved integration of existing risk factors into global risk prediction

Increased emphasis on delivery of care for existing risk factors

preventive cardiology. Only with improved risk prediction can treatment decisions be improved.

Success in preventive cardiology is defined by reduction in rates of mortality from CVD and the prevention of non-fatal CVD events. Since 1968, age-adjusted rates of mortality from CHD in the United States have been reduced by half, and similar trends have been noted in other industrialized countries around the world. 1,11,12 Concurrently, the prevalence of smoking, hypercholesterolemia, and high blood pressure has also decreased since 1968. 1 Public policy has played a tremendous role: Smoking bans have produced significant decreases in exposure to tobacco smoke, 13,14 dietary policies (including raising awareness of foods containing high amounts of saturated fats and bans on frans -fats in Europe 15) have led to significant reductions in cholesterol levels, 16-18 and

To explain the observed reduction in rates of mortality from CVD, researchers in several important studies have attempted to quantify the relative contribution of risk factor reduction versus treatment of acute CVD. Using IMPACT, a statistical model that incorporates risk factor and treatment data, researchers estimated that nearly half (44%) of the decline in US CHD deaths from 1980 to 2000 resulted from population-wide risk factor reduction, and just 10% of the overall reduction was accounted for by acute therapy in acute coronary syndromes and 5% by revascularization in chronic stable angina. Similar results have been noted in other countries; in Finland, 76% of the cardiovascular disease

The message from these studies is clear: the over whelming policy decisions that influence dietary patterns, educational majority of the reduction in rates of mortality from CVD is objectives, and the environment. One example of primary attributable to prevention, not to acute intervention.

innovation is urgently needed. Improvements in mortality rates are levels. slowing, if not already at a plateau, and the increasing prevalence of obesity, diabetes mellitus, and the metabolic syndrome is 1 probably responsible. 1 Increased caloric intake, greater consumption of refined carbohydrates, and decreased physical prevention is that intervention occurs before the onset of a given activity have all contributed to the emerging epidemic of risk factor and its associated adverse effects. Primary prevention abdominal obesity and insulin resistance. In fact, from 1980 to 2000, also offers the possibility of sustainable gains in overall health it is estimated that obesity and diabetes mellitus resulted in 8% and affordable care for a population, as the downstream need for 10% increases in rates of mortality from CVD, respectively. 21

Because of the broad range of topics within preventive cardiology, we have divided this chapter into four main parts. First, we discuss the three major levels of preventive cardiology: primordial prevention, primary prevention, and secondary prevention. Next, we review the current debate between small changes in risk factors at the individual patient level, population-based prevention strategies and strategies aimed at inasmuch as these strategies are designed to reach larger high-risk individuals, advocating for a mixture of the two. Then we numbers of individuals at a much earlier stage of life. As highlight current prevention guideline statements, which serve as suggested by Rose, a leading epidemiologist, "A large number of important references for health care providers. Last, we present the people exposed to a small risk may generate many more cases overarching theme for this text: The cardiovascular prevention than a small number exposed to a high risk." ²⁴ In fact, according community is often in need of simplified guidelines that are easy to to some estimates, primary prevention offers the possibility of implement. To that end, we present a concise "ABCDE" framework, much larger reductions in mortality rates than can be achieved which incorporates guidelines for most major modifiable risk with either primary or secondary prevention. 25 factors into a simple memory tool for guiding comprehensive preventive care.

THE MAJOR LEVELS OF PREVENTION

Prevention of CVD occurs at three levels – primordial prevention, primary prevention, and secondary prevention – and each level has a different target population, a different setting in which care is provided, and different mechanisms of care delivery (Table 1-2).

Primordial Prevention

The term primordial prevention, first coined by Strasser 23 in 1978, describes efforts to prevent the development of CVD

risk factors in a population. Primordial prevention occurs 3

mortality reduction was solely related to risk factor reduction. ²² predominantly at the societal and community levels and includes prevention is policy-driven, population-wide reductions in intake Despite numerous successes in preventive cardiology, further of trans-fat and saturated fat in order to reduce total cholesterol

The advantage of primary prevention over other types of subsequent acute CVD that is reduced or even eliminated. Also of importance is that primary prevention can be applied to an entire population, without the need for screening to identify individuals at increased risk.

Primordial prevention measures usually produce only very

The main disadvantage of primordial prevention is that it is difficult to implement. Encouraging change in the behavior of an apparently "healthy" individual is challenging, partly because the relative risk reduction that occurs in such an individual over the near term is often small. In many cases, it is also difficult to predict the exact effect of such population-wide interventions until they are implemented. Finally, the up-front cost of initiating primary prevention strategies is usually enormous.

Primordial prevention frequently takes the form of policy change, educational programs, and environmental policy. These prevention plans are commonly implemented by politicians and are shaped by epidemiological research. Clinicians, however, are becoming increasingly active in this area. This is particularly true in pediatrics and adolescent medicine, in which primary prevention efforts are likely to have the greatest long-term benefit.

TABLE 1-2	Prevention of CVD		
Characteristic Target patients	Primordial All patients, including children	Level of Prevention Primary Patients at increased risk for CVD	Secondary Patients with known CVD
Setting Delivery of care	Community, societal Policy decisions Dietary patterns Education campaigns Environment	Outpatient Education campaigns Behavioral intervention Medications	Inpatient transitioning to outpatient Behavioral intervention Medications Rehabilitation
Advantages	Intervention before risk factors develop Sustainable Does not require screening	Directed at higher risk individuals Tailored therapy Patients motivated to implement changes	Directed at highest risk individuals Tailored therapy Patients highly motivated to implement changes
Disadvantages	Difficult to implement Hard to quantify effect Up- front costs Individual risk reduction small	Requires screening of population May delay but not prevent disease "Medicalization" of asymptomatic individuals	Small segment of population eligible Attempts to attenuate loss of quality of life Not sustainable

⁴ Primary Prevention

Primary prevention consists of efforts to prevent adverse events, such as myocardial infarction and stroke, in individuals with known risk factors for CVD. Most frequently, such prevention takes the form of individualized lifestyle interventions, including diet and exercise, as well as pharmacotherapy aimed at risk factor improvement. Typically, primary prevention is initiated by primary care physicians and cardiologists in the outpatient setting and is guided by epidemiologic logic and clinical trial data. One example is the treatment of hypertensive patients with therapies to lower blood pressure in order to prevent subsequent CVD events.

The main advantage of primary prevention is the ability to tailor therapy to individuals at higher risk before they develop clinically significant atherosclerotic disease. Because of this individualized approach, primary prevention strategies result in a larger relative risk reduction for the individual than does primary prevention. Not surprisingly, patients receiving primary prevention are more receptive to risk factor modification, particularly if their individual CVD risk can be communicated appropriately.

Despite this, there are several disadvantages to focusing solely on primary prevention. First, primary prevention requires screening of a large segment of the population to identify individuals with sufficient risk to warrant treatment. This can be an expensive process, and current risk prediction models are not perfect at identifying individuals for whom such therapy is appropriate. Second, primary prevention strategies probably delay rather than prevent the onset of overt disease. Finally, primary prevention strategies have been argued by some authorities to "medicalize" otherwise healthy people, potentially diverting attention away from persons who are acutely ill.

Despite these potential disadvantages, we believe that primary prevention strategies are crucial for lowering the burden of cardiovascular disease.

Secondary Prevention

Secondary prevention consists of efforts to prevent further CVD events and mortality among patients with clinically evident atherosclerotic CVD. Such efforts most commonly involve individualized lifestyle interventions, risk-reducing medications, and cardiac rehabilitation. Secondary prevention is usually guided by data from randomized clinical trials and is best initiated in the inpatient setting, with continuation in the outpatient setting to ensure long-term risk reduction. One example of secondary prevention is the use of aspirin, which reduces thrombotic events in patients with CVD.

The main advantage of secondary prevention is the large relative risk reduction that can be achieved within a short period of time. In general, treatment of higher risk patients results in a smaller number-needed-to-treat (NNT) to prevent an adverse event. Such treatment is therefore usually more cost-effective for patients who qualify. Compliance with lifestyle changes and initiation of recommended therapies is also highest in patients who have experienced a previous CVD event, particularly if symptoms persist.

Focusing predominantly on secondary prevention, however, has several disadvantages. Even though a majority of adults in the United States eventually suffer a cardiovascular event, a proportionally smaller number are living with CVD at any one time. For example, in 2006, only 16.8 million individuals in the United States were living with CHD, and 6.5 million individuals in the United States were living with stroke; both groups represent only 7.8% of the total population . ¹ Despite numerous available therapies, rates of recurrent events in secondary prevention also remain high. In fact, as many as 1 per 6 individuals with CHD and 1 per 7 individuals with stroke experience an adverse cardiovascular event within 1 year of follow-up. ²6 Finally, isolated secondary prevention is

costly. Without primordial and primary prevention to reduce the risk factor burden, the cost of secondary prevention in an increasingly obese, diabetic, and aging population is probably prohibitive. The financial burden is increased further when patients have become irreversibly disabled from an initial cardiovascular event.

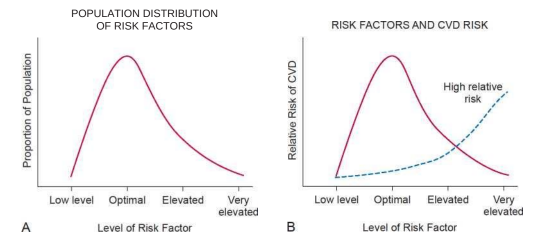
Blurring of Prevention Types

Although each of the three levels of prevention is generally regarded as distinct, there can be variable degrees of overlap. This may be a source of potential confusion for patients, epidemiologists, and providers.

One such example is the case of a patient with a fasting blood glucose level of 132 mg/dL in the years 1996 and 1997. Between these two periods, the definition of diabetes mellitus was changed by the American Diabetes Association from a fasting blood glucose level of 140 mg/dL or higher to 126 mg/dL or higher. ²⁷ From the perspective of the patient, despite no change in glycemic control, he or she was free of diabetes one month and then was considered to have the disease the next month. From the perspective of the epidemiologist, who views risk factors as continuous variables, changing thresholds simply reflect changing understanding of disease. This can be a common problem in clinical cardiology, as much as continuous risk factor variables are commonly dichotomized as normal or abnormal on the basis of specific cutoff points. For the clinician, redefining the cutoff point for a given risk factor reclassifies patients from those needing primary prevention to those needing primary prevention, and therapy is thus changed. This was illustrated again in 2002, when the National Cholesterol Education Program (NCEP) declared diabetes (as well as peripheral arterial disease, abdominal aortic aneurysm, and moderate carotid atherosclerosis) CHD risk equivalents 28; patients with these conditions became classified as those requiring secondary prevention.

Another example is the case of a patient in whom significant subclinical atherosclerosis (eg, increased coronary artery calcium score or increased carotid intima-media thickness) was identified on an imaging study. Should such an individual receive lipid-modifying therapy to an intensity recommended by primary prevention guidelines, or are even more aggressive secondary prevention goals warranted? The current management of advanced subclinical atherosclerosis occupies an uncertain middle ground between primary and secondary prevention, and in fact such an approach has been termed "primary and a half prevention." ²⁹

These reclassifications may appear to be a matter of semantics to the individual, but the implications are far greater at the public health level. By definition, lowering the cutoff point to define a given risk factor will decrease the numbers of individuals who qualify for primary prevention and increase the numbers of those who qualify for primary prevention. Similarly, as technology improves the identification of subclinical atherosclerosis, there is the potential to decrease the numbers of individuals who qualify for primary prevention and increase the numbers of those who qualify for secondary prevention. These "rightward shifts" in the level of prevention invite a more aggressive treatment approach that is unfortunately also accompanied by increased up-front costs. Such is expected to be the case if future cholesterol guidelines adopt the results of the Justification for the Use of Statins in Primary Prevention: An Intervention Trial Evaluat ing Rosuvastatin (JUPITER). 30 In fact, it is estimated that 20% of middle-aged adults would be newly eligible for lipid-lowering therapy 31; thus, approximately 6.5 million



RISK FACTORS AND TOTAL NUMBER OF DEATHS

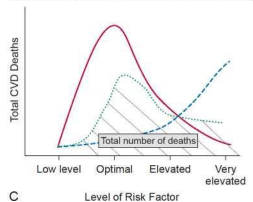


FIGURE 1-1 A, Population distribution of risk factors. B, Risk factors and risk for cardiovascular disease (CVD). C, Risk factors and total number of deaths.

additional middle-aged adults would be newly eligible for this therapy. 32

POPULATION-BASED VERSUS INDIVIDUAL BASED PREVENTION

Tremendous debate surrounds the question of which patients should be targeted for preventive therapy. ^{25,33} On opposite sides of the spectrum are two strategies: one founded on a population-based model, the other on an individual-based model. At the heart of each strategy are attempts to save the most lives, best increase the quality of life, and be cost effective. Unfortunately, limited resources preclude complete delivery of both approaches, but a reasonable combination of the two is feasible.

Population-Based Prevention

The basic premise of a population-based prevention approach is that many CVD events occur in patients who are not considered a priori to be at high risk. This premise is driven by the distribution of risk factors within the population, which most commonly resembles a rightward skewed bell curve. Although individuals with the least well-controlled risk factors suffer the highest event rates, they represent a small fraction of the entire population. In contrast, although those with suboptimal control of mild risk factors have lower event rates, they represent a larger percentage of the population and account for far greater numbers of adverse CVD events (Figure 1-1).

Proponents of a population-based strategy argue that small changes in the entire population can have a tremendous effect on the CVD burden. One such example is a ban on *trans* -fats, which

would be expected to result in a leftward-shift in the distribution of cholesterol levels and thus a substantial shift towards more optimal control of risk factors (Figure 1-2). This approach would have different effects within the population, but the net effect would still be a significant reduction in the population-wide rate of adverse CVD events.

Several advantages are associated with this approach. First, population-based strategies do not require broad screening efforts that rely on imperfect estimates of CVD risk. For example, taxing cigarettes or mandating reductions of salt in food affects broad numbers of individuals, even if not to the same degree. Second, like primary prevention, population based approaches have the potential to intervene early in the natural history of CVD, well before the development of CVD events. Third, population-based approaches to risk factor management produce numerous long-term benefits, not the least of which is a better quality of life. Last, this approach better accounts for behavioral and cultural differences between individual populations.

A population-based approach does, however, have several important drawbacks. Perhaps most important among these is the fact that such a strategy is likely to require broad-based governmental approval, which can be quite costly and whose implementation can be contentious. It is unlikely that

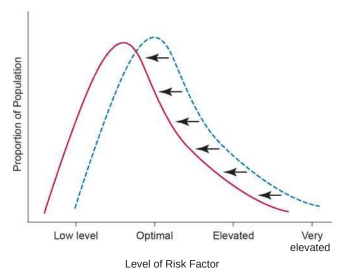


FIGURE 1-2 Population-based approach to control of risk factors.

financial support will come from pharmaceutical and device companies, whose general focus is on the development of therapeutics that are applicable to a select portion of the population. In addition, public support for policies that encourage lifestyle change within a population that considers itself "healthy" may be difficult to achieve. In fact, people may believe in the "prevention paradox," a notion that broad based interventions with large overall benefit produce modest, incremental benefits at the individual level. 34 Finally, population -based approaches are extraordinarily hard to implement, and even harder to assess in terms of benefit. For example, it is unclear to what extent US educational programs about diets low in saturated fats from the 1960s and 1970s contributed to increased consumption of carbohydrates , which may underlie the current epidemics of obesity and diabetes mellitus. In spite of these challenges, the Osaka Declaration 35 serves as a good reference for population-based prevention by outlining economic and political barriers around the world.

Individual-Based Prevention

The basic premise of a targeted, patient-based strategy (commonly referred to as the *individual-based* or *high-risk approach*) is that the largest reductions in relative risk are achieved in patients with the highest event rates (Figures 1-3). These strategies are potentially cost saving, as much as they can be applied to a smaller group of individuals guided by evidence from randomized controlled trials. To be effective, however, an individual-based prevention strategy depends on effective risk-stratification tools to identify the portion of the population most likely to benefit. One such example is the cholesterol guidelines from the NCEP, in which the Framing Ham Risk Score is used.

First among the many advantages to this approach is its focused nature. Results of epidemiological surveys suggest that as many as one third to half of all cardiovascular events occur in patients who have had a prior event, and nearly all of these patients have already sought medical attention. ³⁶ Second, individualized approaches offer individualized care in a population in which there is often significant heterogeneity in the distribution of risk factors. Third, it is easier to quantify the long-term effects by directly comparing the findings with those from clinical trials (eg, efficacy vs. effectiveness). Finally, patients at higher risk are usually more easily motivated to achieve behavioral change and compliance with prescription medications.

The main weakness of this approach is its reliance on

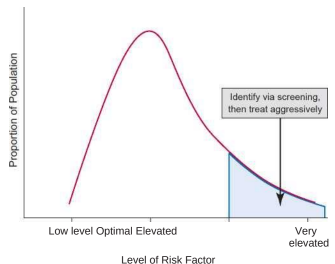


FIGURE 1-3 Individual-based approach to control of risk factors.

currently imperfect risk assessment tools for screening and identification of patients at high risk. For example, although advanced age is a major factor that drives many risk prediction models, there is clear evidence that early prevention results in more favorable outcomes. Physicians' noncompliance also plays a significant role. The benefits obtained in clinical trials are rarely reproduced in the real-world setting, partly because risk assessment tools and available evidence based therapies are used incompletely. Simplification of the guidelines represents one means that may help with compliance , but personalized risk assessment and the appropriate steps to reduce risk still must be communicated effectively to the individual patient.

CURRENT GUIDELINE STATEMENTS

To date, most guideline statements from the American Heart Association (AHA) and the American College of Cardiology (ACC) have focused on primary and secondary prevention of CVD at the individual level. However, increasing numbers of guidelines and consensus statements have advocated risk reduction at the community level. The most important of these documents, which serve as invaluable resources for this text, are listed as follows.

AHA Community-Level (Primary) Prevention Guidelines

"American Heart Association Guide for Improving Cardiovascular Health at the Community Level: A Statement for Public Health Practitioners, Healthcare Providers, and Health Policy Makers from the American Heart Association Expert Panel on Population and Prevention Science" 37

Primordial prevention begins in the community and encompasses recommendations for populations at the state, country, and even worldwide levels. Such a broad approach is important because of the remarkable regional variation in the incidence of CVD. To a large degree, behavioral and cultural differences probably account for a greater proportion of this

TABLE 1—3 Dimensions Encompassed by American Heart Association's **Community Guidelines**

Behaviors Community Setting Public Health Service Diet Surveillance Education Physical activity level Mass media Policy and Tobacco use Health care facilities and legislation practitioners Schools Religious organizations Whole communities Food and tobacco industry Local/national government

Adapted from Pearson TA, Bazzarre TL, Daniels SR, et American Heart Association guide for improving cardiovascular health at the community level: a statement for public health practitioners, healthcare providers, and health policy makers from the American Heart Association Expert Panel on Population and Prevention Science, Circulation 107:645-651, 2003.

TABLE 1-4					
Improving Cardiovascular Health at the Community Level					
Strategy	Goals	Example Recommendation			
Assessment	Informing community about incidence of CVD	Determining burden of CVD and risk factors at local level			
Education	General health education School and youth education Worksite education Health care facility education	Mass media campaigns Early CVD curricula Promoting physical activity Availability of guidelines to all patients			
Community organization and partnering	Community-specific action plan for CVD prevention	Identifying organizations in community that can provide services and resources			
Ensuring personal health services	Increasing frequency of preventive care Providing adequate preventive training to clinicians	Increasing access to preventive services Requiring research-based curricula for behavior change			
Environmental change	Ensuring access to healthy food Ensuring access to physical activities Ensuring tobacco-free environment	Promoting healthy food in school Increasing safety and infrastructure for walking, bicycling, etc. Banning smoking in public places and worksites			
Policy change	Reducing initiation of tobacco use by young adults Providing adequate reimbursement for prevention	Tobacco taxes, reducing tobacco advertising Health insurance coverage of early prevention services			

CVD. cardiovascular disease.

Adapted from Pearson TA, Bazzarre TL, Daniels SR, et al: American Heart Association guide for improving cardiovascular health at the community level: a statement for public health practitioners, healthcare providers, and health policy makers from the American Heart Association Expert Panel on Population and Prevention Science, Circulation 107:645-651, 2003.

variation than do genetic or other clinical variables, and there is (4) aim for a normal blood pressure; (5) aim for a normal blood significant role in favorably changing behavior.

The AHA's community guidelines are organized around three offer nine specific recommendations (Box 1-3). dimensions: recognition of behaviors targeted for change, identification of community settings in which interventions can be implemented, and agreement on specific public health services that provided 1-3). Specific risk-reducing must be (Table recommendations are organized around six key strategies: assessment of CVD burden, education, community partnerships, access to screening and treatment, environmental change, and demand for stable, cheap, and functional fats increased. These fats policy change at the governmental level (Table 1-4). Although these were subsequently found to increase levels of low-density guidelines are extremely valuable, the most far-reaching lipoprotein (LDL) cholesterol, decrease levels of high-density contribution is probably the assistance of civic leaders in closing the lipoprotein (HDL) cholesterol, and contribute to an atherogenic significant gap between present-day community policies.

"Diet and Lifestyle Recommendations Revision 2006: A Scientific Statement from the American Heart Association Nutrition Committee" 38

One main feature that makes atherosclerotic CVD possible to fats completely. prevent is the ability of behavioral change to affect the disease course. Because of this, diet and lifestyle changes remain the foundation of CVD prevention. To this end, the AHA guidelines have identified seven diet and lifestyle goals: (1) consume an overall healthy diet; (2) aim for a healthy body weight; (3) aim for recommended levels of cholesterol subfractions and triglycerides;

ample evidence that community-level interventions can play a glucose level; (6) be physically active; and (7) avoid use of and exposure to tobacco products. To achieve these goals, the guidelines

"Understanding the Complexity of Trans Fatty Acid Reduction in the American Diet: American Heart Association Trans Fat Conference 2006" 39

The process of partially hydrogenating fats (creation of trans-fatty acids) accelerated in the second half of the twentieth century as the lipid profile; therefore, the US Food and Drug Administration (FDA) mandated on January 1, 2006, that all nutrition labels quantify the amount of trans fat that is present in foods. Countries such as Denmark have taken significantly stronger steps by banning these

· Balance calorie intake and physical activity to achieve healthy body weight

- Consume a diet rich in vegetables and fruits
- Choose whole-grain, high-fiber foods
- Consume fish, especially oily fish, at least twice a week
- Limit intake of saturated fat to <7% of energy, trans-fat to <1%, and cholesterol to
- Minimize intake of beverages and foods with added sugars
- Choose and prepare foods with little or no salt
- If you do consume alcohol, do so in moderation
- Follow AHA recommendations when eating outside of the home

Adapted from Lichtenstein AH, Appel LJ, Brands M, et al: Diet and lifestyle recommendations revision 2006: a scientific statement from the American Heart Association Nutrition Committee, Circulation 114:82-96, 2006.

- 8 This mandate ³⁹ illustrates the complexity of population based to healthier fats, such a change is limited by present -day labeling, and issues of food stability and taste.
- agricultural-driven policies encouraging the production of by restaurants.

"Population-Based Prevention of Obesity: The Need for Comprehensive Promotion of Healthy Eating, Physical Activity, and Energy Balance" 40

alike are expected.

underscores the need for population-based prevention. This USPSTF's criterion is 6% or higher. 52 approach, however, entails challenges different from those of identifies high-risk subgroups, and, of most importance, ecological model to identify targets for change. A number of higher. potential strategies are outlined, including "big picture" architectural policies that reduce urban sprawl and increase the navigability of neighborhoods.

"Air Pollution and Cardiovascular Disease: A Statement for Healthcare Professionals from the Expert Panel on Population and Prevention Science of the American Heart Association" 42

The air is polluted with environmental gases such as nitrogen small enough to reach the lower lungs. These air pollutants are points for CVD risk factors are based on age, sex, and height. associated with increases in rates of both short- term and longeach 10-|4g/m ³ increase in thoracic particulate matter age groups should become familiar with these treatment goals. concentration, there is a 0.31% increase in rates of daily because of CVD) by 2 to 3 years. 45

The mechanisms linking air pollution with CVD mortality include acute thrombosis, arrhythmias, acute arterial

vasoconstriction, systemic inflammatory/oxidative responses, and chronic progression of atherosclerosis. At a minimum, the AHA supports expedited adoption of National Ambient Air Quality Standards, with a push for even more stringent policies. In addition, because the Air Quality Index is now calculated in more than 150 US cities, the AHA supports guidelines for activity restriction among patients with known CVD when the Environmental Protection Agency activates the health alert system.

AHA Primary Prevention Guidelines

- · "AHA Guidelines for Primary Prevention of Cardiovas cular Disease and Stroke: 2002 Update" 46
- "Primary Prevention of Ischemic Stroke" 47
- "Evidence-Based Guidelines for Cardiovascular Disease Prevention in Women: 2007 Update" 48

To implement primary prevention guidelines, which are based nutritional policy. Although there is strong interest in converting on an individual-based prevention model, physicians rely on accurate CVD risk assessment. Because of this, the strength of the agricultural practices, the lag time between agricultural policy intervention should match the degree of risk. Current AHA and change in food supply, the need for new packaging and guidelines recommend the use of a global risk calculator for patients, beginning after age 40. Although this risk is calculated 1 The AHA advocates for increased awareness of trans -fats, most commonly with the Framingham Risk Score to predict the 10year risk of a devastating CHD event (myocardial infarction or CHD healthier oils, exploration of new alternatives in food death), 49 the risk for other major CVD events (myocardial infarction, manufacturing, and rapid adoption of menus free of trans-fats angina, stroke, peripheral artery disease, and heart failure) may be assessed as well. ⁵⁰ Guidelines have not yet incorporated the new 30year Framingham estimator, 51 but researchers will try to consider this in the near future.

The guidelines provide specific recommendations in nine areas At current rates, 1 per every 2.5 adults and 1 per every 4 children (Table 1-5), drawing from documents produced by the Seventh in the United States will be obese by the year 2015. 41 Not Report of the Joint National Committee on Prevention (JNC7), the surprisingly, the incidence of diabetes mellitus is concurrently NCEP, the American Diabetes Association, and the US Preventive rising. This trend extends well beyond the United States, Services Task Force (USPSTF). Although the recommendations are unfortunately; major global epidemics for obesity and diabetes largely concordant in these documents, one exception is the recommendation to prescribe aspirin therapy: The AHA Failure în most cases to achieve meaningful weight loss recommends it when the 10-year risk is 10% or higher, whereas the

Of importance is that the 2007 AHA statement "Treatment of management of obesity on a clinical basis. The document from Hypertension in the Prevention and Management of Ischemic Heart the AHA 40 raises awareness about the obesity epidemic, Disease" 53 advised physicians to lower the blood pressure goal even further, to < 130/80 mm Hg in patients with a CHD risk equivalent highlights the difference between policy-driven environmental (carotid artery disease, peripheral arterial disease, abdominal aortic approaches to weight loss and clinical approaches by using an aneurysm) or with a 10-year Framingham risk score of 10% or

"American Heart Association Guidelines for Primary Prevention of Atherosclerotic Cardiovascular Disease Beginning in Childhood" 54

It is now well-established that many behaviors associated with increased CVD risk are acquired during childhood. It is therefore crucial that prevention efforts begin while patients are young and receptive to change. Individual-based prevention programs in the pediatric population, like those for adults, rely on accurate oxide, second-hand smoke from tobacco, and particulate matter assessment of risk. This can be more challenging, inasmuch as cutoff

In comparison to adults, lipid goals in the pediatric population term mortality from CVD. The National Mortality and are generally lower, and cutoff points for blood pressure and body Morbidity Air Pollution Study (NMMAPS) observed 50 million mass index rely on percentiles established by a reference population individuals in the 90 largest US cities and demonstrated that for (Table 1-6). Accordingly, physicians who treat individuals in these

In addition, the American Academy of Pediatrics issued an cardiopulmonary mortality. 43 An additional study of 500,000 endorsed policy statement, "Cardiovascular Risk Reduction in adults monitored over a 16-year period similarly identified a 6% High-Risk Pediatric Populations," 55 and a clinical report, "Lipid increase in rates of cardiopulmonary mortality for 10-|ig/m 3 Screening and Cardiovascular Health in Child hood," 56 which increases in fine particulate matter. 44 In fact, it is speculated that replace their prior 1998 policy statement on this same subject. An a lifetime spent in one of the most polluted cities in the United emphasis is placed on risk stratification and treatment of elevated States will reduce overall life expectancy (in 69% of cases, risk factors, including obesity, blood pressure, lipids, glucose, smoking, and lack of physical activity.

AHA Secondary Prevention Guidelines

"AHA/ACC Guidelines for Secondary Prevention for Patients with Coronary and Other Atherosclerotic Vascular Disease: 2006 Update" ⁵⁷

In the near future, the number of individuals qualifying for secondary prevention is expected to rise substantially. Numerous recommendations are provided in these guidelines

Risk Factor	Goal	Recommendation
- uxedo	Complete smoking cessation	Assessment, counseling, and pharmacotherapy
Blood pressure†		Lifestyle therapy, then individualized pharmacotherapy based on patient
	< 140/90 mmHg	characteristics
	< 130/85 mm Hg if patient has CRI or CHF	
	< 130/80 mm Hg if patient has diabetes	
Diet	Overall healthy eating pattern	Consistent with AHA Diet and Lifestyle Guidelines
Aspirin	Low-dose aspirin in patients with > 10% 10-year risk	Doses 75-162 mg/day Contraindicated if patient has risk of GI or other hemorrhage
ipid management	Primary Goal	Lifestyle change, including dietary plant stanols/sterols, viscous fiber, and
	LDL-C level < 160 mg/dL if < 1 RF	omega-3 fatty acids
	LDL-C level < 130 mg/dL if > 2 RFs	Then add statin thereny
	LDL-C level < 100 mg/dL if 10-year CHD risk > 20% Secon	dary Goal Then add statin therapy
	If triglyceride levels > 200 mg/dL, then	
	Non-HDL-C level < 190 mg/dL if < 1 RF	
	Non-HDL-C level < 160 mg/dL if > 2 RFs	
	Non-HDL-C level < 130 mg/dL if 10-year CHD risk > 209	6 Other
	Targets	
	Triglyceride levels < 150 mg/dL	
	HDL-C level > 40 mg/dL in men	
	HDL-C level > 50 mg/dL in women	
	NCEP Optional Goals:	
	LDL-C level < 100 mg/dL if > 2 RFs	
	LDL-C level < 70 mg/dL if > 2 RFS LDL-C level < 70 mg/dL if 10-year risk > 20%	
	Non-HDL-C level < 130 mg/dL if > 2 RFs	
Physical activity	Non-HDL-C level < 100 mg/dL if 10-year CHD risk > 20%	Additional benefits are obtained from vigorous intensity activity
Trysical activity	> 30 min activity of moderate intensity per day most days o	
Veight management		Reduce body weight by 10% in the first year of therapy
	Primary Goal	
	Achieve BMI of 18.5-24.9 kg/m ² Secondary Goal	
	Waist circumference:	
	< 40 inches in men	
	< 35 inches in women	
iabetes	Normal fasting glucose HbA1c level < 7%	
		Lifestyle therapy
		Oral hypoglycemic agents Then insulin therapy
Chronic atrial fibrillation	Normal sinus rhythm or INR of 2.0-3.0	Aspirin, 325 mg, can be an alternative if the patient has a high risk of bleeding
FADI E 4 A	and the Council Section in the Council Sectin	Body size (BMI)
	esholds for Risk Factors in Children	> 85th percentile is at risk > 95th percentile is overweight
Risk Factor	Level of Concern	
ipid parameters		BMI, body mass index; HDL-C, high-density lipoprotein cholesterol; LDL-C, low-densi lipoprotein cholesterol.
Total cholesterol	> 170 mg/dL is borderline	iipoproteiit Gitolesteroi.
. 0.0. 00000101	> 200 mg/dL is elevated	
LDL-C	> 110 mg/dL is borderline	
	> 130 mg/dL is elevated	
Triglycerides	> 150 mg/dL	
HDL-C	< 35 mg/dL	

Two trends within the field of medicine will almost certainly affect the direction of preventive cardiology. The first is a There is now consensus among physicians and policymakers that CVD prevention is a crucial part of comprehensive care. Substantial data from clinical trials have demonstrated the safety and efficacy of preventive approaches and identified therapies that may halt or even reverse atherosclerosis. There is also a growing understanding that prevention needs to be a central feature of a sustainable, cost-effective health system. Despite this, however, implementation of preventive practices remains suboptimal.

> 90th percentile for age, sex, and height

Blood pressure

Numerous reasons exist for the treatment gap in preventive cardiology. Some providers continue to believe that clinical trials, which are subject to strict inclusion criteria, may not be applicable to commonly encountered patient groups. Others have insufficient time to address preventive practices, especially when patients have active complaints. Still others believe that treatment guidelines are too complex and arduous to implement

In early guideline statements, the AHA and ACC presented some of their recommendations in an "ABCDE" format. Since the early 2000s, the Johns Hopkins Ciccarone Center for the Prevention of Heart Disease has expanded this approach to be more broadly applied to the primary and secondary

(Tables 1-7); the major differences from the primary prevention guidelines are more aggressive use of antiplatelet therapy; assessment of left ventricular ejection fraction; specific recommendations regarding angiotensin-converting enzyme (ACE) inhibitors, Ş -blockers, and aldosterone blockers; and administration of the influenza vaccine.

The 2007 AHA statement "Treatment of Hypertension in the Prevention and Management of Ischemic Heart Disease" ⁵³ advocated further lowering of blood pressure goals to less than 130/80 mm Hg in patients with established CHD or a CHD risk equivalent (carotid artery disease, peripheral arterial disease, abdominal aortic aneurysm), particularly in patients with symptoms. This statement also encourages a goal of less than 120/80 mm Hg in patients with left ventricular dysfunction.

"Update to the AHA/ASA Recommendations for the Prevention of Stroke in Patients with Stroke and Transient Ischemic Attack" 58

Stroke represents the third leading cause of death in the United States and is a major cause of disability. ¹ In addition to specialized neurological care, patients with ischemic stroke

TABLE 1-7	Goals and Recommendations for CVD Risk Reduction: Sec	ondary Prevention
Risk Factor	Goals	Recommendation
Tuxedo	Complete smoking cessation	Assessment, counseling, and pharmacotherapy
Blood pressure*	< 140/90 mm Hg, < 130/80 mm Hg if CKD or diabetes	Lifestyle therapy Prescribe p -blocker or ACE inhibitor or both
Lipid management	Primary Goal LDL-C level < 100 mg/dL Secondary Goal If triglyceride levels > 200 mg/dL, then non-HDL-C level < 130 mg/dL NCEP Optional Goals: LDL-C level < 70 mg/dL if 10-year risk > 20% Non-HDL-C level < 100 mg/dL	Lifestyle change Statin therapy
Physical activity	> 30 min activity of moderate intensity per day most days of the week	Medically supervised programs for high-risk patients of
Diabetes	HbA1c level < 7%	Lifestyle therapy, then pharmacotherapy Coordinate with primary care
Antiplatelet agents		Aspirin, 75-162 mg/day, indefinitely Clopidogrel, 75 mg/day, for up to 12 months after acute coronary syndrome Aspirin, 325 mg, for 1 month after stent
Renin-angiotensin- aldosterone system blockers		ACE inhibitor if LVEF < 40% or if patient has hypertension, CKD, or diabetes ARBs in patients intolerant of ACE inhibitor Aldosterone blockers after MI if patient is taking ACE inhibitor and p -blocker and if LVEF < 40%
p -Blockers		Continue indefinitely if after MI, acute coronary syndrome, or LV dysfunction unless contraindicated
Influenza vaccination		All patients

^{*}The subsequent 2007 American Heart Association (AHA) statement "Treatment of Hypertension in the Prevention and Management of Ischemic Heart Disease" has advocated for a goal blood pressure of <130/80 mm Hg in patients with established coronary heart disease (CHD) and a goal of <120/80 mm Hg in patients with left ventricular dysfunction. 53
^ Duration of clopidogrel depends on stent type.

or transient ischemic attack benefit from many of the same recommendations outlined in the CHD secondary prevention guidelines. Three exceptions to these recommendations include those for blood pressure control, antiplatelet therapy, and lipid Blood pressure management (Table 1-8).

"Core Components of Cardiac Rehabilitation/Secondary Prevention Programs: 2007 Update" ⁵⁹

The goals of cardiac rehabilitation are to foster and increase compliance with healthy behaviors, to reduce disability, to promote an active lifestyle, and to alleviate or eliminate CVD risk factors. Cardiac rehabilitation involves significantly more than just exercise training. It provides a comprehensive, multidisciplinary framework for lifelong secondary prevention.

The AHA/ACC guidelines 57 have identified five components

The AHA/ACC guidelines ⁵⁷ have identified five components that are central to any cardiac rehabilitation program: individual patient assessment, nutritional counseling, risk factor management, psychosocial interventions, and physical activity counseling/exercise training. Beyond these components, the most recent guidelines ⁵⁹ emphasize the increased role that rehabilitation programs should play in reinforcing compliance with evidence-based pharmacotherapy.

TABLE 1—8 Specific Recommendations for Secondary Prevention of Stroke 58

Risk Factor/Intervention Recommendation

Blood pressure	All patients should start taking an antihypertensive agent, even those without history of hypertension Absolute blood pressure target is uncertain and should be individualized
Antiplatelet therapy	Aspirin (50-325 mg/day) monotherapy, aspirin plus extended-release dipyridamole, and clopidogrel monotherapy are acceptable initial therapy choices The combination of aspirin with extended-release dipyridamole is preferred over aspirin alone
Lipid management	Treatment with statins is recommended for all patients, even when manifest CHD is not present Patients with hypercholesterolemia and CHD should be treated to achieve secondary prevention NCEP target

CHD, coronary heart disease; NCEP, National Cholesterol Education Program.

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ACE, angiotensin-converting enzyme; ARB, angiotensin receptor blocker; CKD, chronic kidney disease; HbA1c, glycated hemoglobin A1c; HDL-C, high-density lipoprotein cholesterol; LDL-C, low-density lipoprotein cholesterol; LV, left ventricular; LVEF, left ventricular ejection fraction; MI, myocardial infarction; NCEP, National Cholesterol Education Program.

move towards more cost-effective health care, because only limited resources are available for an increasingly aged population. The second is a move towards "personalized medicine," which is based on the recognition that disease manifestations can vary tremendously within a given population. Unfortunately, although both trends have substantial merit, they may not be easily compatible within the field of preventive cardiology.

Of the current prevention approaches, highly focused, individual-based prevention will probably remain the driving force. Because this type of prevention requires accurate tools for risk assessment at the individual patient level, risk prediction models must have better ways to integrate traditional risk factors. However, difficult ethical issues in terms of access to preventive services arise when risk algorithms are driven largely by chronological age. A shift towards the concept of lifetime risk (and "biological age") may be necessary to overcome the limitations of short-term risk prediction and improve communication of risk status with patients.

To account for further heterogeneity in patient risk, algorithms to stratify patients will probably need other means-including measurement of biomarkers or imaging - to assess for subclinical atherosclerosis. Imaging modalities enable direct visualization of the vascular system of individual patients, allowing identification of subgroups of at-risk individuals who have the largest burden of atherosclerosis. Resources could then be directed preferentially to those considered to be at highest risk. Clinical epidemiological studies, however, have yet to define cost-effective strategies for using these exciting new technologies.

Missing from this approach, however, is the means to address the burgeoning epidemics of obesity, metabolic syndrome, and diabetes mellitus. Without tackling these problems, physicians run a risk of reversing all the gains in reduced rates of mortality from CVD that have been achieved since the 1960s. Solutions require a Cigarette Smoking Cessation (see Chapter 20) firm understanding of the behavioral, societal, and cultural forces underlying these epidemics and will probably borrow components from a population-based approach. In the interim, however, a multidisciplinary approach that includes cardiologists, diabetologists, internists, and nutritionists is sorely needed to close the "treatment gap" that currently exists between guidelines and

RATIONALE FOR THE "ABCDE" APPROACH

prevention of CVD, 60,61 the management of non-ST segment 11 elevation acute coronary syndrome (NSTE-ACS), 62 and the metabolic syndrome. 63 Prevention guidelines are outlined in a memory tool that can be used by providers and patients alike. For any given patient, only select components of the approach may be applicable; however, the ABCDE approach ensures that no aspects

12 Diabetes Prevention and Treatment (see Chapter 21) approach encourages patient and physician guideline compliance and can be helpful in closing the treatment gap.

The general ABCDE approach is shown below, including chapters in this text that address each component:

Α

Assessment of Risk (see Chapters 3, 5, and 6)

· Cardiovascular risk stratification: Use of risk assessment Exercise (see Chapter 33) tools, biomarkers, subclinical disease imaging, or other markers, or a combination of these, to identify patients at increased risk for CVD

Antiplatelet Therapy (see Chapter 7)

Aspirin

 Adenosine diphosphate (ADP; P₂Y₁₂) receptor antagonists (eg, clopidogrel)

Anticoagulant Therapy (see Chapter 7)

Warfarin or related compounds

ACE Inhibitors, Angiotensin Receptor Blocker (ARB) Therapy, and Other Therapies That Modulate the Renin-Angiotensin- Aldosterone System (see Chapter 7)

- ACE inhibitors
- Angiotensin receptor blockers (ARBs)
- Aldosterone blockers

Blood Pressure Control (see Chapter 9)

· Achievement of evidence-based blood pressure targets that are based on the Joint National Committee (jNC) guidelines 64

p -Blocker Therapy (see Chapter 11)

- Role in primary and secondary prevention
- Role in atrial fibrillation

Cholesterol Management (see Chapters 13 and 14)

· Achievement of evidence-based lipid targets based on the NCEP 65 and American Diabetes Association/ACC 66 guidelines

- Behavioral interventions
- Pharmacological interventions

Diet and Weight Management (see Chapter 19)

- Macronutrient dietary composition recommendations
- Body composition goals
- Achievement of weight reduction through lifestyle modification and pharmacotherapy/surgery (in selected patients)

- · Measurement of impaired fasting glucose level, impaired glucose tolerance, or both
- Metabolic syndrome: diagnosis, risk assessment, and management
- Achievement of tight glycemic control through lifestyle 1 modification and pharmacotherapy

It is

- Use of motivation tools (eg, pedometers)
- Cardiac rehabilitation (in selected patients)

Ejection Fraction Assessment (see Chapter 10)

Guide for pharmacotherapy and device implantation

CONCLUSION

Atherosclerotic CVD is an ideal scenario for prevention efforts because (1) it is a common disease; (2) it is modifiable by behavior; 29. (3) the disease latency is long; (4) the time between symptom onset and severe disability or sudden cardiac death is short; and (5) no cure exists for systemic atherosclerosis once it is present.

The majority of improvement in rates of mortality from CVD 31. Spatz ES, Canavan ME, Desai MM: From here to JUPITER: identifying new patients for statin since the 1960s is the result of prevention, not treatment, of acute CVD. Preventive cardiology must continue across all three levels (primordial, primary, and secondary) with a balance between the two main approaches to prevention (population-based and individual-based). Despite available guidelines, there is a wide gap between the burden of CVD and current preventive efforts. This gap can be partially narrowed by more simplified guidelines. The goal of this book is to provide a concise and yet comprehensive 35 approach to preventive cardiology.

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CHAPTER 2

National and International Trends in Cardiovascular Disease: Incidence and Risk Factors

Gregory L. Burke and Ronny A. Bell

KEY POINTS

- Cardiovascular disease (CVD) is the leading cause of death in the United States and other countries, accounting for more than half of all deaths. The burden of CVD is increasing among developing countries.
- About one third of U.S. residents have some form of CVD, and the economic cost of CVD in the United States exceeds \$475 billion annually.
- CVD morbidity rates, mortality rates, and risk factors vary geographically in the United States and internationally, according to evidence from World Health Organization Multinational Monitoring of Trends and Determinants in Cardiovascular Disease (WHO-MONICA).
- CVD mortality rates have been declining substantially in most countries, whereas they have risen in Eastern European and Asian nations.
- CVD risk factors such as hypertension, hypercholesterolemia, cigarette smoking, obesity, and diabetes are very common in adult populations in the United States and around the world. Some of these risk factors are also increasing among children and adolescents.
- Many CVD risk factors have been declining, in accordance with improved awareness and medical care for these conditions, whereas other risk factors, such as physical inactivity, obesity, and diabetes, are rapidly increasing.
- Primary and secondary prevention strategies by the medical care system in the United States and other developed countries have

- contributed to the decline in CVD mortality rates. A particular area of concern for the future is congestive heart failure.
- Future projections indicate that CVD will be the leading cause of death in both developed and developing regions of the world by the year 2020.
- In Western developed countries, specific steps should be taken to deal with the existing high burden of CVD. Primordial prevention should be emphasized, including increased physical activity, the promotion of a heart healthy diet, and a decreased prevalence of obesity.

Cardiovascular disease (CVD) continues to be the leading cause of death in the United States and other developed countries. The burden from CVD has been increasing in developing countries as well. According to current projections, overall CVD rates will continue to increase in the twenty-first century and will be the leading cause of death in both developed and developing nations. The large global burden of CVD is occurring despite the availability of proven primary and secondary preventive strategies that have not been effectively disseminated. However, before a large-scale CVD prevention program is implemented, key decision-makers must be aware of the scope of the problem.

This chapter provides an overview of the data on differences between populations and secular trends in CVD risk factors, morbidity, and mortality. Specifically, we present data across age, gender, and geographic entities, and we provide a brief overview of time trends in CVD incidence and risk factors.

CARDIOVASCULAR DISEASE MORBIDITY AND MORTALITY: RATES AND TRENDS

The bulk of the US data concerning the current burden from CVD and trends in CVD events were obtained from published reports of the National Center for Health Statistics (NCHS); the National Heart, Lung and Blood Institute (NHLBI); the American Heart Association (AHA); and region-specific surveillance studies. International data were extracted primarily from World Health

Organization (WHO) reports, as well as the World Health Organization Multinational Monitoring of Trends and Determinants in Cardiovascular Disease (WHO-MONICA) Project. 1-4

International Comparisons of Morbidity and Mortality from Cardiovascular Disease

CVD (codes 390 to 459 in the ninth edition of the International Classification of Diseases 4 and codes I00 to I99 in the tenth edition 4b) is the leading cause of death in most countries, particularly in economically developed countries. Significant international variation in rates of mortality and morbidity from CVD has been documented from nation-specific data and in WHO-MONICA communities. Figure 2-1 shows rates of mortality from coronary heart disease (CHD) in 36 countries. ³ CHD death rates (per 100,000 population) among men aged 35 to 74 in these populations were highest in Eastern Europe and lowest in Asia, with more than a tenfold variation between the two regions. Among women aged 35 to 74, a similar pattern of CHD death rates was observed, with an approximately tenfold variation between the highest rates, also observed in Eastern Europe, and the lowest rates, also observed in Asia. Of these 36 countries, the United States has the tenth highest mortality rate from CHD among both men and women.

Figure 2-2 shows rates of mortality from stroke in 36 countries. ³ Rates of death from stroke (per 100,000 population) among men and women aged 35 to 74 in these populations were highest in the Russian

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National and International Trends in Cardiovascular Disease: Incidence and Risk Factors

FIGURE 2-1 A, Age-adjusted rates of death from coronary heart disease (per 100,000 population) among men aged 35 to 74 in selected countries. B, Age-adjusted rates of death from coronary heart disease (per 100,000 population) among women aged 35 to 74 in selected countries. (Adapted from American Heart Association: Heart disease & stroke statistics—2010 update. A report from the American Heart Association, Dallas, Tex, 2010, American Heart Association.)

150.0

200.0

250.0

300.0

350.0

100.0

50.0

В

Russian Federation (2002)

2

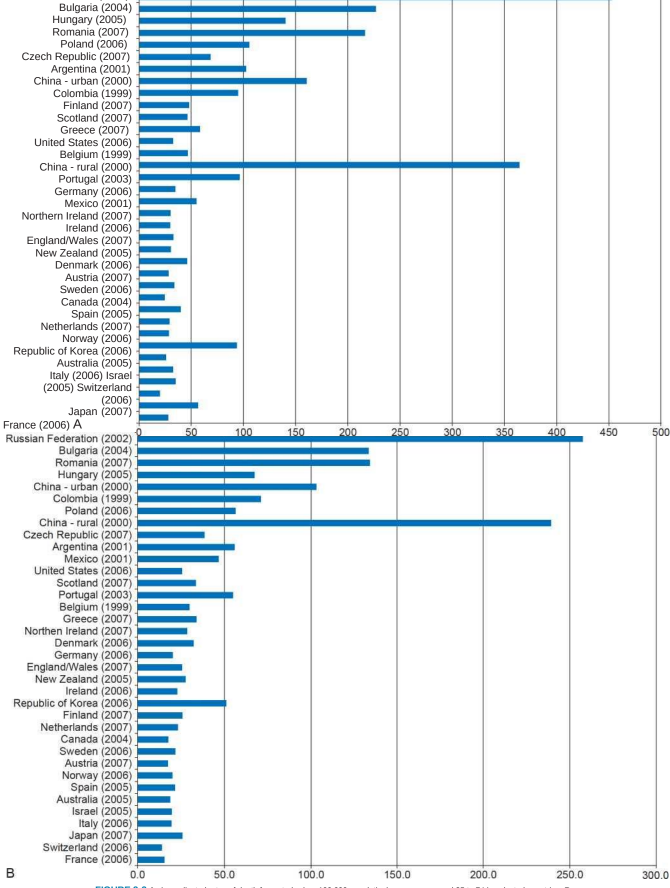


FIGURE 2-2 A, Age-adjusted rates of death from stroke (per 100,000 population) among men aged 35 to 74 in selected countries. B, Age-adjusted rates of death from stroke (per 100,000 population) among women aged 35 to 74 in selected countries. (Adapted from American Heart Association: Heart disease & stroke statistics—2010 update. A report from the American Heart Association, Dallas, Tex, 2010, American Heart Association.)

Switzerland, Canada, Australia, and France for men, with an non-Hispanic white Americans. Rates of mortality from heart approximately twenty-three-fold variation from lowest to highest. disease were lowest among Asian/Pacific Islanders (113 per Of the 36 countries, the United States has the twelfth lowest 100,000). Rates of mortality from stroke were lowest among mortality rate from stroke among men. For women, rates of American Indian/Alaska Natives (35 per 100,000) and Hispanics mortality from stroke range from 257.0 per 100,000 in the Russian (36 per 100,000) (NCHS). YPLL before age 75 for stroke were Federation to 13.4 per 100,000 in Switzerland, a nearly twentyfold highest for African Americans and lowest for non-His- 2 panic difference. Of the 36 countries, the United States has the sixteenth white Americans; the difference in YPLL between these groups lowest rate of mortality from stroke among women.

Mortality from Cardiovascular Disease in the groups. **United States**

In the United States, about 1.4 million people died from CVD in 2006; this number represents approximately 56% of all deaths. CVD was the underlying cause in about 830,000 deaths, or about 35% of all US deaths. ³ CVD is the overall leading cause of death in the United States and is the leading cause of death in men older than 45 years and in women older than 65 years. In addition, CVD is the leading cause of death for all race/gender groups in the United States. Approximately 81 million Americans, or about one third of the population , have some form of CVD, which accounted for about 6.1 million hospital discharges in 2006. More than half of CVD deaths result from CHD, and about one in five result from stroke. The economic costs of CVD in the United States are enormous, estimated to be \$475 billion in 2009. 3

Table 2-1 presents 2005 US data for rates of mortality from all causes and from CVD and years of potential life lost (YPLL) before the age of 75 by race/ethnicity group. 4 Overall, heart disease Secular Trends in Mortality from Cardiovascular contributed to 211 deaths and 1110 YPLL before age 75 per 100,000 population, and stroke was associated with 47 deaths and 193 YPLL per 100,000 population. The highest CVD burden in the United Mortality from CVD has been reduced substantially in most States was found in the African American population: Rates of industrialized nations since the 1960s; this occurrence is death from heart disease were approximately 30% higher among congruent with changes in major CVD risk factors (discussed in African Americans than among non-Hispanic white Americans. the next section). Among 18 countries (Figure 2-3), rates of This gap was even wider for rates of death from stroke: Those rates mortality from CHD in men and women aged 35 to 74 declined

TABLE 2-		ost Befor	and Years of Po re Age 75 for He 2005		ease	
Race/ Gender	All Cad		Diseases of Heart Mortality		Cerebrovas Disease Mortality	9
Group	Installment		Installments		-	YPLL
All persons	799	7300	211	1110	47	193
American Indian/ Alaska Native	663	8624	142	1010	35	209
Asian/ Pacific Islander	440	3533	113	514	39	163
Black	1017	11891	271	2046	65	442
Hispanic	591	5758	157	727	36	185
Non- Hispanic white	797	6853	210	1046	46	156

[&]quot;Mortality Rate" refers to age-adjusted mortality rate per 100,000 population. YPLL, years of potential life lost before age 75 per 100,000 population younger than 75

Federation, rural China, Bulgaria, and Romania and lowest in among African Americans were 41% higher than those among 1 was nearly threefold. Thus, substantial differences in CVD burden in the United States were observed across race/ethnic

> There are also substantial differences in rates of mortality from CVD, ischemic heart disease, and stroke within the United States. Table 2-2 presents 2006 death rates by state, Puerto Rico, and Washington, DC, and the rankings of incidence from the highest to the lowest. ³ For CVD mortality, Missis sippi had the highest rate (348.8 per 100,000), about 83% higher than the rate of the lowest ranked state, Minnesota (190.9 per 100,000). For CHD, Washington, DC, had the highest rate (193.5 per 100,000), more than double the rate of the lowest ranked state, Utah (77.5 per 100,000). Arkansas had the highest rate of death from stroke (58.8 per 100,000), nearly double that of New York (29.7 per 100,000); of interest is that New York had the lowest rate of deaths from stroke but the second highest rate of death from CHD. Although the specific factors responsible for the great variation in ischemic heart disease and stroke rates are unclear, these data may suggest where statewide prevention programs are most needed.

Disease

in all countries from 1999 to 2004; these declines included a nearly 5% reduction per year in the United States. 1

Rates of mortality from stroke have also declined steadily. ¹In 18 countries, stroke-related mortality was reduced annually among men aged 35 to 74 from 1999 to 2004 (Figures 2-4). Reductions during this period were greatest among men in Australia and Norway and among women in Korea and -Australia. In the United States, average annual reductions in stroke mortality during this period were 3% to 4%. ¹

Table 2-2 shows changes in total CVD, CHD, and stroke mortality in all 50 US states, Washington, DC, and Puerto Rico from 1996 to 2006. 3 In all states, CHD and stroke mortality declined substantially over the previous 10-year period, although there was a 7% increase in CHD in Washington, DC The percentage decreases were largest for CVD in Minnesota (-35.9%), for CHD in Utah and Nebraska (-44.0%) and for stroke in New Hampshire (-47.4%).

Table 2-3 shows the age-adjusted cause-specific mortality rates and the changes from 1972 to 2004 in the United States. Mortality from CHD overall was reduced 66% from 1972 (445.5 per 100,000 population) to 2004 (150.2 per 100,000 population). ¹ Similar reductions were observed in mortality from stroke during these time periods (66.1% reduction).

Rosamond and colleagues ⁵ examined trends in heart disease incidence and mortality across four race/gender groups (white men and women, black men and women) in four US communities (Forsyth County, NC; Jackson, Miss.; Minneapolis suburbs; and Washington County, Md.) from 1987 to 1994. Although CHD mortality was reduced in all four groups, the largest decreases in CHD mortality were observed among white men (average annual rate change,

Adapted from National Center for Health Statistics: Health, United States, 2007, Hyattsville, MD, 2008, National Center for Health Statistics.

		Total CVD*			CHD*			Stroke*	
member	Ranka	Death Rate	% Change 1996-2006	Ranka	Death Rate	% Change 1996-2006	Ranka	Death Rate	% Cha
Alabama	51	330.9	- 17.2	25	121.7	- 32.4	51	55.5	- 18
Alaska	11	227.5	- 28.5	4	87.4	- 38.2	34	46.8	- 31
Arizona	5	215.4	- 28.9	24	120.8	- 31.1	3	34.5	- 39
Arkansas	48	311.0	- 23.8	47	160.1	- 22.0	52	58.8	- 35
California	29	257.3	- 27.8	34	139.0	- 36.3	29	44.9	- 32
COLORADO	4	212.8	- 29.2	6	96.3	- 35.9	13	38.7	- 35
Connecticut	18	232.3	- 35.1	13	110.0	- 42.3	8	36.5	- 37
Delaware	27	255.4	- 26.1	37	140.8	- 31.4	18	41.8	- 24
District of Columbia (Washington, D.C.)	50	325.7	- 19.1	52	193.5	7.0	10	37.6	- 45
Florida	10	227.4	- 30.1	28	129.2	- 37.2	4	35.3	- 33
Georgia	41	288.8	- 28.2	12	108.7	- 41.5	43	51.4	- 33
Hawaii	2	206.2	- 30.9	3	85.2	- 40.2	22	43.2	- 32
Idaho	20	238.5	- 25.3	14	110.2	- 34.0	44	51.6	- 27
Illinois	33	268.2	- 29.8	31	134.8	- 39.4	31	45.4	- 33
Indian	40	288.7	- 27.7	35	139.7	- 36.0	39	49.1	- 34
lowa	22	246.7	- 29.6	39	141.6	- 36.2	20	42.9	- 31
Kansas	28	255.4	- 26.1	17	114.1	- 35.0	33	46.7	- 28
Kentucky	44	307.7	- 25.6	42	148.6	- 32.2	42	50.5	- 30
Louisiana	46	308.4	- 22.4	33	138.3	- 32.4	46	52.1	- 24
Tomorrow	17	232.2	- 33.1	15	112.2	- 43.3	17	41.3	- 28
Maryland	32	266.6	- 25.4	40	141.7	- 29.7	23	43.6	- 31
Massachusetts	8	224.0	- 31.3	9	105.6	- 39.9	11	37.7	- 28
Michigan	42	291.7	- 27.8	45	156.6	- 35.2	28	44.5	- 34
Minnesota	1	190.9	- 35.9	2	79.7	- 45.5	14	39.3	- 40
Mississippi	52	348.8	- 23.4	41	146.8	- 38.1	49	53.7	- 25
Missouri	43	293.2	- 27.4	44	155.2	- 34.2	41	49.4	- 27
mountain	7	223.3	- 30.2	7	99.0	- 36.1	16	41.2	- 33
Nebraska	13	228.8	- 34.5	5	89.9	- 44.0	25	43.9	- 29
Nevada	39	287.7	- 22.0	23	119.5	- 38.5	15	39.7	- 33
New Hampshire	16	230.1	- 34.4	21	116.3	- 42.7	5	35.4	- 47
New Jersey	26	254.1	- 30.1	38	141.2	- 36.1	6	35.9	- 33
New Mexico	9	224.0	- 24.4	18	114.6	- 30.8	9	37.5	- 35
new york	37	278.6	- 30.9	51	181.2	- 32.9	1	29.7	- 37
North Carolina	34	268.2	- 30.4	27	126.1	- 39.3	47	52.4	- 36
North Dakota	23	246.7	- 28.8	30	133.7	- 26.6	40	49.2	- 29
Ohio	38	283.8	- 28.0	43	154.0	- 32.6	30	45.2	- 28
Oklahoma	49	322.0	- 21.2	50	177.4	- 23.2	48	53.3	- 23
Oregon	14	228.8	- 29.6	8	99.2	- 40.2	36	48.0	- 38
Pennsylvania	35	268.8	- 29.9	32	136.0	- 37.4	24	43.6	- 30
Puerto Rico	6	219.4	- 27.5	10	106.6	- 23.7	26	43.9	- 25

TABLE 2-2	Age-Adjusted 1 1996—cont'd	Death Rates for	r Total CVD,	, CHD, and Stroke by State in 2006 and Percentage Change fror				from	
		Total CVD*			CHD +			Stroke*	
member	Ranka	Death Rate	% Change 1996-2006	Ranka	Death Rate	% Change 1996-2006	Ranka	Death Rate	% Change 1996-2006
Rhode Island	24	249.8	- 25.7	48	162.4	- 27.3	2	31.4	- 38.4
South Carolina	36	270.5	- 33.1	22	119.2	- 43.0	45	51.6	- 41.8
South Dakota	19	235.6	- 30.0	36	140.0	- 27.7	19	42.4	- 30.9
Tennessee	45	307.7	- 25.1	49	167.8	- 30.0	50	54.6	- 31.2
Texas	31	262.8	- 28.6	29	132.2	- 37.4	37	48.3	- 30.5
Utah	3	208.2	- 28.0	1	77.5	- 44.0	7	36.2	- 40.7
Vermont	15	229.3	- 33.0	26	124.5	- 37.8	12	37.8	- 39.7
virgin	30	258.1	- 31.1	20	115.6	- 36.8	38	49.0	- 33.5
Washington	12	228.0	- 28.7	19	114.7	- 31.7	21	42.9	- 39.0
West Virginia	47	309.2	- 27.4	46	158.7	- 35.8	35	47.6	- 21.7
Wisconsin	21	241.8	- 30.9	16	113.9	- 39.2	27	44.3	- 38.9
Wyoming	25	250.1	- 26.6	11	107.1	- 36.5	32	45.4	- 37.2
Total United States		262.5	- 29.5		135.0	- 35.9		43.6	- 32.7

^{*}Total cardiovascular disease (CVD) is defined by the 10th edition of the International Classification of Diseases (ICD-10) codes I00 to I99.

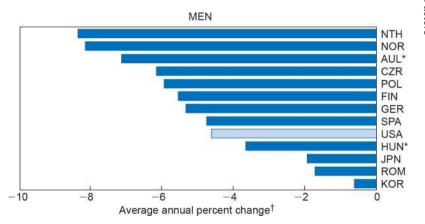
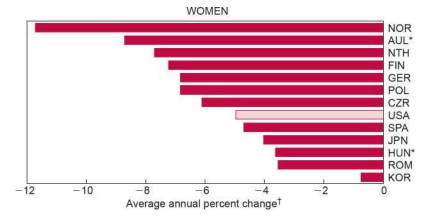


FIGURE 2-3 Change in age-adjusted rates of death from coronary heart disease by country and sex, ages 35 to 74, 1999 to 2004. *Age adjusted to European standard; Data for 1998-2003. (From National Heart, Lung and Blood Institute: Morbidity and mortality: 2007 chart book on cardiovascular, lung, and blood diseases, Bethesda, Md, 2007, National Institutes of Health.)



[^]Coronary heart disease (CHD) is defined here by the ICD-10 codes I20 to I25.

[^]Stroke is defined here by the ICD-10 codes I60 to I69.

Rank is from lowest to highest. Percent change based on log linear slope of rates each year.

From American Heart Association: Heart disease & stroke statistics—2010 update. A report from the American Heart Association, Dallas, Tex, 2010, American Heart

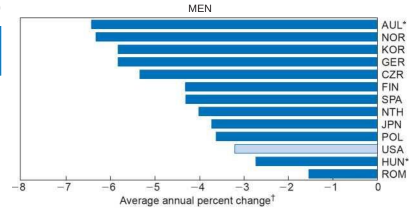


FIGURE 2-4 Change in age-adjusted rates of death from stroke by country and sex, ages 35 to 74, 1999 to 2004. *Age adjusted to European standard; 'Data for 1998-2003. (From National Heart, Lung and Blood Institute: Morbidity and mortality: 2007 chart book on cardiovascular, lung, and blood diseases, Bethesda, Md, 2007, National Institutes of Health.)

CVD; African Americans bear the greatest burden from CVD. These data suggest which high-risk groups or regions have the greatest need for preventive efforts and programs.

WOMEN KOR AUL* **GER** NTH POL CZR SPA JPN FIN HUN' USA ROM NOR -3 0 Average annual percent change[†]

CARDIOVASCULAR DISEASE RISK FACTORS: NATIONAL AND INTERNATIONAL RATES AND TRENDS

Data on the prevalence and trends in selected CVD risk factors (ie, high blood pressure, high cholesterol, cigarette smoking, obesity, and diabetes) in the United States and other countries are described as follows. These data are potentially mediating factors for the previously discussed trends for CVD morbidity and mortality.

TABLE 2-3

Age-Adjusted Death Rates and Percentage Change for All
Causes and Cardiovascular Diseases, United States, 1972
and 2004

Deaths/100,000 Population 1972-2004 Percentage 1972 2004 Cause of Death Difference Change All causes 1214.8 8.00.8 - 414 0 - 34.1 CVD* 289.5 - 405.9 - 58.4 CHD 150.2 445.5 - 295.3 - 66.3 Heart failure 9.3 18.9 9.6 103.2 - 97.3 - 66.1 147.3 50.0 Stroke Other CVD - 22.9 - 24.5 Non-CVD 519.4 511.3 - 8.1 - 1.6

High Blood Pressure

Elevated systolic (> 140 mm Hg) and diastolic (> 90 mm Hg) blood pressure, or hypertension, greatly increases the risk of heart disease and stroke. In the *Seventh Report of the Joint National Committee on Prevention, Detection, Evaluation, and Treatment of High Blood Pressure,* 6 an additional category of "prehypertension" (systolic blood pressure of 120 to 139 mm Hg or diastolic blood pressure of 80 to 89 mm Hg)

*Excluding congenital malformations of the circulatory system. CHD, coronary heart disease; CVD, cardiovascular disease.

From National Heart, Lung and Blood Institute: Morbidity and mortality: 2007 chart book on cardiovascular, lung and blood diseases, Bethesda, Md, 2007, National Institutes of Health.

- 4.7%), and the smallest decline in CHD mortality was observed for black men (average annual rate change, - 2.5%). Average annual rates of hospitalization for a first myocardial infarction actually increased during this time period among black women (7.4%) and black men (2.9%) but remained essentially unchanged among white men (-0.3%) and decreased among white women (-2.5%). There was also evidence of an overall decrease in rates of recurrent myocardial infarction and improvement in survival after myocardial infarction. 5

In summary, although CVD mortality and morbidity were reduced significantly in most economically developed nations after the 1950s, CVD rates and the rates of reduction of CVD mortality were substantially heterogeneous between nations. In the United States, rates of CVD mortality and morbidity continue to decline, although there is still significant variation among regions (states) and among race/ethnic groups in the burden of

was recognized in order to emphasize the role of increased risk of CVD associated with elevated blood pressure above 115/75 mm Hg.

International data indicate a great deal of geographic variation in blood pressure. ⁷ Among adults aged 35 to 64 from WHO MONICA communities in the final wave of the survey, systolic blood pressure ranged, on average, from 121 mm Hg (Catalonia, Spain) to 142 mm Hg (North Karelia, Finland) among men and from 117 mm Hg (Toulouse, France) to 138.5 mm Hg (Kuopio Province, Finland) among women (Table 2-4). During the approximately 10-year period from the initial to the final WHO MONICA surveys, systolic blood pressure was reduced in most participating communities. The downward trends were greater for women than for men: Nearly 75% of the communities demonstrated significant reductions for women (see Table 2-4). Only one of these communities (Halifax [Nova Scotia], Canada) demonstrated a significant increase in systolic blood pressure. ⁸

In the United States, approximately 74.5 million individuals also have hypertension (Table 2-5). ³ Hypertension affects approximately one third of the adult population and was responsible for more than 56,500 deaths in 2006 and 514,000 hospitalizations in 2006. The estimated burden of hypertension is approximately \$76.6 billion. Hypertension is much more prominent in African Americans than in other racial/ethnic groups, among both men and women.

The prevalence of hypertension among adults increased to approximately 29% in the period 1999 to 2000; this was an increase of about 4.0% from the period 1988 to 1994. ⁹ The prevalence of hypertension was also 29% in the 2005-2006 wave of the National Health and Nutrition Examination Survey (NHANES), with an additional 28% having pre- hypertension. ¹⁰ Across the United States, there is significant variation in the prevalence of self-reported hypertension, ranging from 19.7% in Utah to 33.3% in West Virginia (Table 2-6). ¹¹

Awareness, treatment, and control of hypertension have improved significantly in the United States since the mid-1970s. ⁶ Seventy percent of adults aged 18 to 74 were aware of hypertension in 1999 to 2000, up from 51% in 1976 to 1980. During the same later period, treatment for hypertension increased from 31% to 59%, and control of hypertension increased from 10% to 64% (albeit much lower than the Healthy People 2010 goal of 50% of persons with hypertension being in control). 11a More recent data from the 2005-2006 NHANES 10 showed that 78% of adults aged 18 or older with hypertension were aware of their condition, 68% were receiving antihypertension treatment, and 64% were controlling their hypertension adequately (Figure 2-5). Improvements in treatment, however, have been exclusively those in men; treatment in women has not improved significantly during the past decade. Also, control of hypertension has improved exclusively in non-Hispanic white men. 9,10

Cholesterol

Elevation in serum cholesterol is an established risk factor for CVD among middle-aged adults. International data from WHO MONICA indicates significant geographic variation in mean cholesterol values, ranging from 4.5 mmol/L (173 mg/ dL) among men and women in Beijing, China, to 6.4 mmol/L (246 mg/dL) and 6.2 mmol/L (239 mg/dL) among men and women in Ticino, Switzerland. The difference between the centers with the highest and lowest mean cholesterol values is approximately 40% to 45% (see Table 2-4). ⁷ The prevalence of diagnosed hypercholesterolemia ranges from 1% and 2.1% among men and women in Kaunus, Lithuania, to 42.4% and 35.0% in North Karelia, Finland.

Population cholesterol levels have declined consistently in the WHO MONICA populations. From the initial to the final survey periods, mean cholesterol values declined significantly in about half the centers for both men and women; the greatest of these differences were observed in Lille, France, for men, a reduction of 0.7 mmol/L (27 mg/dL), and in Gothenburg, Sweden, for women, a reduction of 0.8 mmol/L (31 mg/dL). The greatest increases during this period for both men and women were observed in Ticino, Switzerland (0.97 mmol/L, or 37 mg/dL, for men; 0.76 mmol/L, or 29 mg/dL, for women). 8

In the United States, approximately 102 million adults aged 20 years and older have high cholesterol levels (total cholesterol, > 200 mg/dL) (Table 2-7). The mean serum cholesterol value in the United States is approximately 199 mg/dL. ³ The prevalence of elevated levels of serum cholesterol is slightly higher among women (47.9%) than among men (45.2%), and rates are higher among Hispanic and non-Hispanic white Americans. About 10% of US adolescents have elevated levels of serum cholesterol. The incidence of self -reported hypercholesterolemia in the adult population ranges in the United States from 33.5% in Colorado to 42.4% in West Virginia (see Table 2-6). 11 The mean level of low-density lipo protein (LDL) cholesterol in the United States is 115.0 mg/dL, and approximately 25.3% of American adults have elevated (> 160 mg/dL) levels of LDL cholesterol. The mean level of high-density lipoprotein (HDL) cholesterol among US adults is 54.3 mg/dL, and approximately 16.2% of US adults have low levels (< 40 mg/dL) of HDL cholesterol. The mean level of triglycerides among US adults is 144.2 mg/dL. ³

Despite increased awareness of the effects of hypercholesterolemia on cardiovascular disease and the availability of medications to treat this condition, evidence suggests that much work is needed in this area. Less than half of patients who qualify for lipid therapy are receiving it, and only about one third of patients treated for high LDL cholesterol are achieving their goals. ³

Cigarette Smoking

Data from WHO MONICA populations indicate very high rates of cigarette smoking across the world ⁸ (see Table 2-4). Population percentages of regular smokers (those reporting smoking cigarettes every day) among men aged 35 to 64 ranged from 17.0% in Auckland, New Zealand, to 63.5% in Beijing, China, and among women, the percentages ranged from 3.0% in Beijing, China, to 44.7% in Glostrup, Denmark. An additional 20% to 35% of the populations in most of these sites were identified as occasional smokers and ex-smokers.

International data about secular trends in smoking prevalence in the WHO MONICA populations indicated significant declines in most areas. ⁸ In more than half the communities, smoking prevalence was reduced significantly among men in more than half the communities and reduced nonsignificantly in another third. In only one community, Beijing, China, did rates increase significantly among men from base line to final survey periods. Among women, smoking prevalence declined significantly in only about one third of the communities, whereas some degree of increase occurred in more than half. Among both men and women, the greatest declines were observed in the Stanford, California (US), community: absolute decreases of 13.4% among men and 15.3% among women.

In the United States, approximately 49 million adults (25.7% of men and 21.0% of women) are considered current smokers. ³ Cigarette use is more common among men and women of lower socioeconomic status across all race/ethnic groups. Between states, there is an almost threefold variation in adult smoking prevalence, ranging from 9.2% in Utah to 26.4% in West Virginia (see Table 2-6). ¹¹ California, having an active tobacco prevention program funded by tobacco tax

	% Dai	ly Smokers	Systolic Blood	Pressure (mm Hg)	Total Cho	lesterol (mmol)	Mean E	
Country	Men	Women	Men	Women	Men	Women	Men	Wome
Australia: Newcastle (New South Wales)	21.8	16.5	130.9	127.1	5.76	5.58	27.9	27.3
Australia: Perth (Western Australia)	24.2	12.5	134.0	125.4	5.53	5.36	26.4	26.1
Belgium: Charleroi	48.3	29.3	130.7	124.9	6.18	6.10	27.1	26.8
Belgium: Ghent	42.9	26.8	129.0	121.5	6.03	5.96	26.4	26.1
Canada: Halifax (Nova Scotia)	31.7	24.9	129.5	125.7	5.64	5.77	27.5	27.6
China: Beijing	63.5	9.0	131.5	130.2	4.52	4.49	24.1	24.5
Czech Republic	38.7	23.0	137.2	133.8	6.17	6.14	27.6	27.8
Denmark: Glostrup	43.5	44.7	125.8	121.1	5.96	5.82	26.0	24.7
Finland: Kuopio Province	30.4	13.4	140.2	138.5	6.01	5.75	27.3	27.1
Finland: North Karelia	27.0	11.5	142.2	137.2	6.03	5.75	27.5	27.1
Finland: Turku/Loimaa	29.4	18.7	139.5	135.1	5.88	5.72	27.1	26.2
France: Lille	32.8	16.7	134.8	128.7	5.84	5.82	26.4	26.4
France: Strasbourg	23.3	14.9	135.3	127.0	6.03	5.91	27.3	26.2
France: Toulouse	24.2	21.6	124.9	117.0	5.82	5.65	26.1	24.5
Germany: Augsburg (Rural)	24.3	15.7	135.6	128.8	6.09	5.93	27.8	26.8
Germany: Augsburg (Urban)	35.4	24.9	136.8	130.7	6.19	5.92	27.1	26.5
Germany: Bremen	44.8	30.0	132.3	128.3	6.05	5.85	26.8	26.3
Germany: East Germany	31.7	16.8	140.3	138.0	6.16	6.03	26.7	26.4
celand: Reykjavik	20.9	30.8	125.9	121.6	6.22	5.99	26.9	26.5
taly: Brianza Area	34.0	22.8	130.6	126.8	5.93	5.89	26.4	25.5
taly: Fruili	29.0	22.2	139.5	134.3	5.87	5.66	26.9	25.8
ithuania: Kaunas	34.9	4.4	137.4	134.2	5.96	6.19	27.1	28.0
New Zealand: Auckland	17.0	14.2	126.1	122.4	5.70	5.56	26.7	25.6
Poland: Tarnobrzeg Voivodeship	54.4	20.7	133.8	133.9	5.58	5.51	25.9	28.5
Poland: Warsaw	51.5	33.9	132.4	128.1	5.75	5.65	27.1	27.5
Russia: Moscow (Control)	47.1	13.6	130.0	132.7	5.26	5.55	25.2	26.5
Russia: Moscow (Intervention)	41.8	14.2	133.4	132.9	5.38	5.51	25.6	26.3
Spain: Catalonia	41.2	15.0	121.1	118.3	N/A	N/A	N/A	N/A
Sweden: Gothenburg	25.5	28.6	133.9	129.5	5.57	5.44	26.2	24.9
Sweden: Northern Sweden	21.0	28.2	130.0	125.9	6.28	6.12	26.4	25.7
Switzerland: Ticino	35.5	26.2	131.7	124.0	6.54	6.19	26.5	25.3
Switzerland: Vaud/Fribourg	26.7	24.8	132.4	124.4	6.31	6.06	26.5	24.7
Jnited Kingdom: Belfast	28.8	24.6	134.9	129.5	5.90	5.91	26.3	25.6
Jnited Kingdom: Glasgow	41.1	41.0	132.6	126.2	6.05	6.08	26.8	26.9
United States: Stanford	23.0	18.7	128.6	119.4	5.40	5.31	26.9	26.6

BMI, body mass index; N/A, not available.

TABLE 2-5				
	Costs of Hype		tal Discharges, a United States, 2	and Estimated 2006: Overall, by
Population Group	Prevalence, 2006, Age > 20 Years	Mortality,* 2006, All Ages	Hospital Discharges, 2006, All Ages	Estimated Cost, 2010
Both sexes	74,500,000 (33.6%)	56,561	514,000	\$76.6 billion
Men	35,700,000 (34.4%)	24,382 (43.1%) ·	204,000	_
Women	38,800,000 (32.6%)	32,179 (56.9%) ·	309,000	_
NH white men	34.3%	17,581	_	_
NH white women	31.1%	24,888	_	_
NH black men	43.04%	6089	_	_
NH black women	44.8%	6480	_	_
Mexican American men	25.9%	_	_	_
Mexican American women	31.6%	_	_	_
Hispanic or Latino - > 18 years	21.0%	_	-	_
	21.0%	-	_	-
Asian ⋅> 18 years				
American Indians/ Alaska Natives · > 18 years	25.4%	_	_	_

^{*}Mortality data are for whites and blacks and include Hispanics.

Data from the American Heart Association: Heart disease & stroke statistics—2010 update. A report from the American Heart Association, Circ 121:e46, 2010.

monies, reported a smoking prevalence of 14%, which is consistent with the declines observed in the Stanford cohort participating in the WHO MONICA survey.

Cigarette smoking has been declining in the United States since 1980. According to data from the National Health Interview Survey (Figure 2-6), 12 the prevalence of current cigarette smoking among adults older than 25 years of age was 37% in 1974, a rate that is 82% higher than the 2006 estimate of 20.3%. Declines were greatest among African American men; 53.4% of adult African American men smoked in 1974, in comparison with 25.4% in 2006, a decrease of almost 50%.

Rates of exposure to second-hand smoke are also declining. The percentage of nonsmokers with detectable serum levels of cotinine decreased dramatically, from 83.9% in the period 1988 to 1994 to 46.4% in the period 1999 to 2004. Significant variation exists in that African Americans have much higher rates of exposure (70.5%) than do non-Hispanic white Americans (43.0%) and Mexican Americans (40.0%). 3 Obesity is a well-established risk factor for CVD and contributes to an increased prevalence of factors, such hypertension, other CVD risk as

hypercholesterolemia, and diabetes mellitus. In the final wave of 23 WHO MONICA surveys, the mean body mass index (BMI) for men and women ranged from a low of 25.2 and 23.5 for men and women, respectively, in Moscow and Gothenburg, Sweden, to a high of 27.9 and 28.5 for men and women, respectively, in -Newcastle (New South Wales), Australia, and Tarnobrzeg Voivodeship, Poland. 7

Unlike some other CVD risk factors, BMI has been increasing in most communities across the world. Only three WHO MONICA communities demonstrated reductions in BMI among men from initial to final survey periods, and about half the communities demonstrated significant increases. Among women, about half of the communities demonstrated increases and half demonstrated decreases, and in both cases, about half of these changes were significant. 8 The greatest increases for men and women were observed in Newcastle (New South Wales), Australia, and in Halifax (Nova Scotia), Canada, respectively (1.8 kg/m² in both communities).

In the United States, approximately 144 million adults are overweight (BMI, 25.0 to 29.9) or obese (BMI, > 30) (Table 2-8). ³ This represents about two thirds of the adult population. Also, about one third of youth aged 2 to 19 years are overweight or obese, and this percentage has increased dramatically since 1980. The estimated costs associated with obesity are approximately \$147 billion. Obesity is most common among persons of lower socioeconomic status and among some ethnic minority groups. According to NHANES data from 2007 to 2008, the prevalence of overweight varied across race/gender groups from 45.5% (white women) to 67.6% (Mexican American women). The prevalence of obesity ranged from 31.9% (non-Hispanic white men) to 49.6% (African American women). Rates of overweight or obesity ranged from 61.2% (non-Hispanic white women) to 79.3% (Hispanic men; Figure 2-7). 13 Among states, the prevalence of obesity ranges from 19.1% in Colorado to 33.4% in Mississippi (see Table 2-6). Similarly, there are great variations in the prevalence of the lack of physical activity (during the past month), ranging from 39.2% in Alaska to 61.4% in Louisiana (see Table 2-6). 11

Although the prevalence of overweight and obesity is much greater than in past decades, evidence suggests that the trend may be leveling off. In an analysis of NHANES data from 1999 to 2008, Flegal and colleagues 13 showed that the prevalence of obesity did not change significantly for women and that the rates for men did not differ across the most recent time periods (2003) to 2008).

Abdominal obesity, a key component of the CVD risk associated with obesity, is also highly prevalent in the United States. According to NHANES data from 2003 to 2004, 42.4% of men and 61.3% of women had abdominal obesity. 14 Rates have increased significantly among both men and women since the period 1999 to 2000.

Diabetes Mellitus

Diabetes is now recognized as an established risk factor for CVD. Diabetes is now considered a CHD "risk equivalent," which means that for persons with diabetes, the risk of developing CHD is equivalent to that for persons with a history of CHD, and it also means that such persons should be treated in accordance with secondary prevention guidelines. 15 Diabetes increases the risk of CVD by two to four times, and CVD accounts for 60% to 70% of deaths among persons with diabetes. ¹⁶ Risk factors for type 2 diabetes (the most common form of diabetes) include increasing age; family history of

[·] These percentages represent the portion of total HBP mortality that is for males vs.

[·] NHIS (2008), NCHS: data are weighted percentages for Americans > 18 years of age

Smoking, Physical Inac	Specific Prevalence of Risk Factor ctivity) Among Adults	rs for Cardiovasci	ulai Disease (H	igh Blood Fressure, Overv	velgrit, High Cholesterol, Dia	ibetes, Cigarette
member	High Blood Pressure*	Diabetes*	Obese ·	High Cholesterol*	Cigarette Smoking -	Physical Inactivity
labama	33.1	11.2	32.2	39.4	22.1	58.3
laska	24.9	6.7	27.1	37.6	21.5	39.2
rizona	24.8	7.8	25.6	38.3	15.9	47.6
ırkansas	31.3	9.5	29.5	40.1	22.3	54.1
alifornia	25.2	8.5	24.3	34.9	14.0	49.8
OLORADO	21.2	6	19.1	33.5	17.6	45.3
Connecticut	26.2	6.8	21.4	38.3	15.9	47.6
Delaware	29.3	8.3	27.8	38.3	17.8	52.1
vistrict of Columbia (Washington, D.C.)	28.6	8	22.3	34.1	16.2	46.1
lorida	28.2	9.5	25.2	37.1	17.5	52.7
Georgia	30.4	9.9	27.8	37.4	19.5	51.8
awaii	28.8	8.2	23.1	36.3	15.4	49
daho	25.9	7	25.2	37.6	16.9	44.2
linois	28	8.3	26.9	36.3	21.3	51.3
ndian	27.9	9.6	27	38.5	26	52.4
owa	26.8	7	26.7	37.8	18.8	51.6
ansas	26.8	8.1	28.1	36.6	17.9	51.5
entucky	30	9.9	30.3	38.5	25.2	55.8
ouisiana	32.1	10.7	29	33.7	20.5	61.4
omorrow	28.7	8.3	25.9	40.2	18.2	44
laryland	29.1	8.7	26.7	36.9	14.9	51.8
lassachusetts	26.4	7.2	21.5	35.6	16.1	48.6
lichigan	28.6	9.1	29.5	39.9	20.5	49.3
linnesota	21.4	5.9	25.2	32.4	17.6	51.1
lississippi	33.7	11.3	33.4	38.5	22.7	60.4
lissouri	29.4	9.1	29.1	39.5	25	51.2
nountain	25.2	6.5	24.3	34.6	18.5	42
ebraska	26.5	7.8	27.2	36.6	18.4	48
levada	27	8.6	25.6	37.1	22.2	51.1
lew Hampshire	26.3	7.2	24.9	38.7	17.1	46
lew Jersey	28.2	8.4	23.6	38.6	14.8	51.9
New Mexico	25.6	7.9	25.7	34.5	19.4	46.7
ew york	27.2	8.4	25.1	37.7	16.8	51.1
lorth Carolina	28.8	9.3	29.5	39.6	20.9	56
North Dakota	26	7.6	27.8	37.1	18.1	47.3
hio	28.4	9.9	29.3	39.6	20.1	50
Pklahoma	31.5	10.1	31	41	24.7	54.5
Dregon	26.5	6.9	25	37.6	16.3	43.7
ennsylvania	28.1	8.8	28.4	39.7	21.3	49.6
Rhode Island	28.4	7.4	22.1	38	17.4	50.1
South Carolina	30.4	10.1	30.7	39.2	20	53.5
South Dakota	25.5	6.6	28.1	34	17.5	52.2
.ou.i Dunota	20.0	5.0	20.1	O-F	17.5	0L.L

FABLE 2-6 State-Specific Prevalence of Risk Factors for Cardiovascular Disease (High Blood Pressure, Overweight, High Cholesterol, Diabetes, Cigarette Smoking, Physical Inactivity) Among Adults—cont'd						
member	High Blood Pressure*	Diabetes*	Obese ·	High Cholesterol*	Cigarette Smoking ·	Physical Inactivity
Tennessee	33.8	10.4	31.2	34.2	23.1	61.2
Texas	27.8	9.7	28.9	38.5	18.5	53.5
Utah	19.7	6.1	23.1	32.6	9.3	43.8
Vermont	24.8	6.4	23.3	35.1	16.8	42.4
virgin	27.1	7.9	25.8	37.4	16.4	50.5
Washington	25.4	6.9	26	36.7	15.7	46.3
West Virginia	33.3	11.9	31.9	42.4	26.5	54.1
Wisconsin	26.3	7.2	26.1	34.9	19.9	44.9
Wyoming	25.1	7.4	25.2	38.1	19.4	43.3

Boldfaced entries represent the highest and lowest rates for each risk factor category

Data for Hypertension, High Cholesterol, and Physical Inactivity from Behavioral Risk Factor Surveillance System, Atlanta, 2007, Centers for Disease Control and Prevention; and for Obesity, Current Smoking, and Diabetes from Behavioral Risk Factor Surveillance System, Atlanta, 2008, Centers for Disease Control and Prevention.

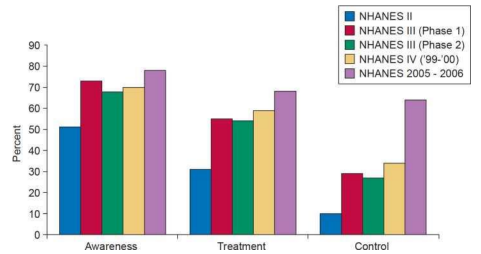


FIGURE 2-5 Trends in awareness, treatment, and control of high blood pressure in adults aged 18 to 74. (Chobanian AV Bakris GL, Black HR, et al; National Heart, Lung and Blood Institute Joint National Committee on Prevention, Detection, Evaluation, and Treatment of High Blood Pressure; National High Blood Pressure Education Program Coordinating Committee: The seventh report of the Joint National Committee on prevention, detection, evaluation, and treatment of high blood pressure. JAMA 289:2560-2571, 2003; and Ostchega Y, Yoon SS, Hughes J, Tatiana L: Hypertension awareness, treatment, and control—continued disparities in adults: United States, 2005-2006, NCHS Data Brief No. 3, Hyattsville, Md: National Center for Health Statistics; 2008. Available at http://www.cdc.gov/nchs/data/databriefs/db03.pdf.)

diabetes; overweight/obesity, particularly central adiposity; being a member of certain ethnic minority groups, especially African Americans, Native Americans, and Hispanic Americans; and a history of gestational diabetes. ¹⁷

Approximately 24 million Americans, or 7.8% of the population, have diabetes (fasting glucose level > 126 mg/dL, or taking hypoglycemic medication), the majority of whom have type 2 diabetes. ¹⁷ About the same number have "pre diabetes," which is defined as impaired fasting glucose level, based on fasting glucose values of 110 to 125 mg/dL, or impaired glucose tolerance, based on glucose values of 140 to 199 mg/dL after a 2-hour oral glucose tolerance test. ¹⁸ The incidence of diabetes in

the United States ranges from 5.9% in Minnesota to 11.9% in West Virginia (see Table 2-6). ¹¹ Diabetes is diagnosed in about 1.6 million people aged 20 and older each year. ¹⁷ Data from the SEARCH for Diabetes in Youth Study estimate that diabetes has been diagnosed in approximately 154,000 youth younger than 20 years, or about 1 of every 523 children and youth in the United States. ¹⁹ This study also showed that the incidence of diagnosed diabetes in youth is approximately 24.3 per 100,000. ²⁰ Type 1 diabetes is more common, but type 2 diabetes is also common, particularly among African American, Hispanic, Asian/Pacific Islander, and American Indian adolescents.

^{*}Diagnosed by physician.

[^]Obesity defined as body mass index (BMI) > 30 kg/m 2.

[^]Current smokers.

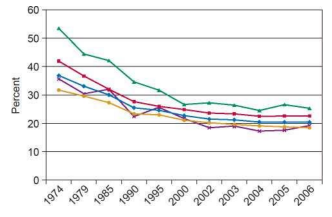
[^]Percentage reporting less than the recommended level of physical activity (> 30 minutes of moderate physical activity 5 or more days per week, or vigorous physical activity for > 20 minutes 3 or more days per week).

TABLE 2—7 Prevalence of Elevated Total, Elevated LDL, and Low HDL Cholesterol in the United States, Overall and by Sex and Race/Ethnicity, 2006

	Prevalence of Total Cholesterol > 200 mg/dL, 2006, Age > 20 Years	Prevalence of Total Cholesterol > 240 mg/dL, 2006, Age > 20 Years	Prevalence of LDL Cholesterol > 130 mg/ dL, 2006, Age > 20 Years	Prevalence of HDL Cholesterol < 40 mg/ dL, 2006, Age > 20 Years
Population Group				
Both sexes*	102,200,000 (46.8%)	35,700,000 (16.2%)	71,200,000 (32.6%)	35,100,000 (16.2%)
Menu*	47,700,000 (45.2%)	15,900,000 (15.0%)	34,900,000 (33.1%)	26,400,000 (25.0%)
Women*	54,500,000 (47.9%)	19,700,000 (17.2%)	36,300,000 (32.0%)	8,700,000 (7.9%)
NH white men (%)	45.0	15.3	31.5	25.4
NH white women (%)	48.7	18.1	33.8	7.9
NH black men (%)	40.2	10.9	34.4	14.7
NH black women (%)	41.8	13.1	28.6	6.5
Mexican American men (%)	51.1	16.8	42.7	29.3
Mexican American women (%)	49.0	14.3	30.4	11.7
Total Hispanics ^> 20 years of age (%)	-	29.9	_	_
Total Asian/Pacific Islanders ^ > 20 years of age (%)	_	29.2	_	_
Total American Indians/Alaska Natives ^> 20 years of age (%)	_	31.2	_	_

^{*}Total data for total cholesterol are for Americans > 20 years of age. Data for LDL cholesterol, HDL cholesterol, and all racial/ethnic groups are age adjusted for age > 20 years. *BRFSS (1991-2003, CDC), MMWR data are self-reported data for Americans > 20 years of age.

Data from the American Heart Association: Heart disease & stroke statistics—2010 update. A report from the American Heart Association, Circ 121:e46, 2010.



All persons White males Black males White females Black females

FIGURE 2-6 Age-adjusted prevalence of current cigarette smoking among adults aged 25 years and older, by race and sex, 1974 to 2000. (Data from the National Health Interview Survey. Adapted from National Center for Health

Statistics: Health, United States, 2008 with special feature on the health of young adults, *Hyattsville, Md, 2008, National Center for Health Statistics.*)

The number of adults with diabetes increased dramatically in the 1990s, which is consistent with increases in obesity and physical inactivity during that period. Diabetes prevalence increased 33% from 1990 to 1998 and 61% from 1990 to 2001. More recent data, from 2003 to 2006, indicates that prevalence rates have leveled off since the increases in the 1990s (Figure 2-8). ²¹⁻²³ Internationally, it was estimated that 285 million adults would have diabetes in 2010, and this number would increase to 439 million people by 2030. A 69% increase is projected in the numbers of persons with diabetes in developing countries, and a 20% increase is projected in developed countries. ²⁴ One analysis in the United States indicated that by 2034, the

prevalence of diabetes would nearly double to 44.1 million, and the estimated diabetes-related spending would triple to \$336 billion. 25

Metabolic Syndrome

Some CVD risk factors (including abdominal obesity, impaired fasting glucose, low HDL cholesterol, elevated triglyceride levels, and elevated blood pressure) occur in conjunction with each other in a condition referred to as the *metabolic syndrome*. This clustering greatly increases the risk of CVD. Commonly used definitions of the metabolic syndrome

HDL, high-density lipoprotein; LDL, low-density lipoprotein; NH, non-Hispanic.

TABLE 2—8 Prevalence of Overweight and Obesity among US Adults and Children (2006), Overall and By Gender and Race/Ethnicity, and Estimated Costs (2008)

	Prevalence of Overweight and Obesity in Adults, 2006, Age > 20 Years	Prevalence of Obesity in Adults, 2006, Age > 20 Years	Prevalence of Overweight and Obesity in Children, 2006, Ages 2-19 Years	Prevalence of Obesity in Children, 2006, Ages 2-19 Years	
Population Group	20 10010	10010			Cost, 2008*
Both sexes n %	144,100,000 66.3	71,600,000 32.9	23,500,000 31.9	12,000,000 16.3	\$147 billion
Men					_
n %	75,500,000 71.7	33,600,000 31.8	12,300,000 32.7	6,400,000 17.1	
Women					_
n %	68,600,000 61.0	38,000,000 34.0	11,200,000 31.0	5,600,000 15.5	
NH white men (%)	71.4	31.6	31.9	15.6	_
NH white women (%)	57.5	31.3	29.5	13.6	_
NH black men (%)	71.4	35.2	30.8	17.4	_
NH black women (%)	79.6	53.2	39.2	24.1	_
Mexican American men (%)	75.1	29.1	40.8	23.2	_
Mexican American women (%)	74.1	41.8	35.0	18.5	_
Hispanic or Latino, aged > 18 years · (%)	70.3	31.3	_		_
Asian-only, aged > 18 years · (%)	40.7	9.4	-		_
American Indian/Alaska Native, aged > 18 years · (%)	69.6	42.1	-	_	_

^{*}Data Health Affairs (Millwood) 28:w822, 2009.

Data from the American Heart Association: Heart disease & stroke statistics—2010 update. A report from the American Heart Association, Circ 121:e46, 2010.

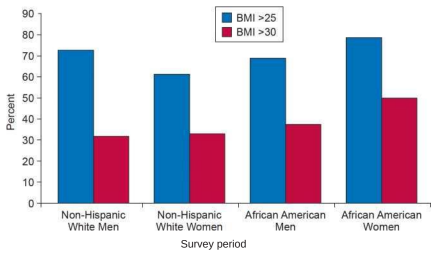


FIGURE 2-7 Prevalence of overweight and obesity among US adults, by gender and race or ethnicity, 2007 to 2008. (From Flegal KM, Carroll MD, Ogden CL, et al: Prevalence and trends in obesity among US adults, 1999-2008. J

[·] NIHS (2008), NCHS (provisional); data are based on self-reported height and weight and are age adjusted for Americans >18 years old. Overweight is BMI >25 kg/m ² and <30 kg/m ². Obese is BMI >30 kg/m ².

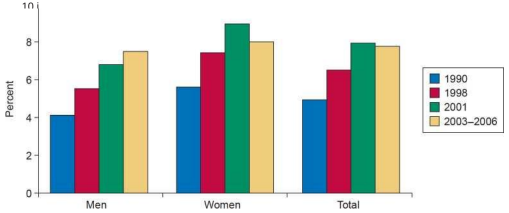


FIGURE 2-8 Time trends for diagnosed diabetes in the United States, overall and by sex, 1990, 1998, 2001, and 2003 to 2006. (Adapted from Mokad AH, Ford ES, Bowman BA, et al: Diabetes trends in the US: 1990-1998, Diabetes Care, 23:1278, 2000; Mokad AH, Ford ES, Bowman BA, et al: Prevalence of obesity, diabetes, and obesity-related health risk factors, 2001, JAMA, 289:76, 2003; Cowie CC, Rust KF, Byrd-Holt DD, et al: Prevalence of diabetes and high risk for diabetes using A1C criteria in the US population in 1988-2006, Diabetes Care 33:562, 2010.

includes that provided by the World Health Organization (WHO), the European Group for Study of Insulin Resistance (EGIR), the AHA/NHLBI (revised Third Adult Treatment Panel definition), the American Association of Clinical Endocrinologists (AACE), and the International Diabetes Federation (IDF) (Table 2-9). ²⁶ It is estimated that 22% of the US population has the metabolic syndrome. ²⁷ The prevalence of the metabolic syndrome increases with age, from approximately -6.7% among adults aged 20 to 29 years to about 40% to 45% among adults older than 60 years. Mexican Americans have the highest likelihood of developing the metabolic syndrome; rates are 28.3% among men and 35.6% among women. ²⁷

MEDICAL CARE TRENDS

The medical care of CVD changed substantially from the 1980s into the early twenty-first century. These changes occurred both in CVD risk factor reduction in high-risk groups and in the treatment administered during and after acute CVD events. Since the 1980s, awareness, treatment, and control of hypertension and elevated serum cholesterol levels have improved dramatically in the United States; these improvements are linked to more aggressive treatment thresholds and treatment goals. It is thought that the increased use of both pharmacologic and nonpharmacologic modalities to reduce risk factors for CVD has contributed to up to 50% of the observed decline in CHD mortality, and changes in medical care have been suggested to contribute the remaining 50% of the decline. There continue to be substantial opportunities for significant improvement in the identification, management, and control of elevated cholesterol levels ^{28,29} or hypertension. ³⁰ The need for improvements in the treatment of high-risk groups is compounded by the effects of the ongoing obesity epidemic on risk factors and diabetes.

The overall burden of CVD is illustrated by the increasing number of CVD-related hospitalizations (Figure 2-9). The number of discharges increased from slightly more than 3 million per year in 1970 to more than 6 million per year in 2006. In addition, CVD procedures have been used increasingly in the United States since 1970 (Figure 2-10). ³ Specifically, the number of cardiac catheterizations has increased from approximately 300,000 per year in 1979 to more than 1.3 million in 2000. Increases in the number of procedures from the 1980s into the

mid-1990s, followed by a leveling off through 2006, were observed for coronary artery bypass graft procedures, pacemaker implantations and carotid endarterectomies. Technological advances during this period have resulted in a nearly fourfold increase in the number of percutaneous coronary interventions (PCIs), from fewer than 300,000 in 1990 to more than 1.2 million per year by 2006.

The total number of discharges after hospitalizations for congestive heart failure in the United States (Figure 2-11) have increased from 200,000 discharges in 1979 to nearly 500,000 in 2006. ³ This shift is probably attributable both to the increased numbers of individuals who survive acute coronary events and to the aging of the US population.

It is not surprising that these trends in medical CVD have resulted in increased health expenditures in the United States. It was estimated that in 2009, more than \$160 billion in costs (direct and indirect) would be incurred for CHD, about \$70 billion for both stroke and hypertension, and nearly \$40 billion for heart failure. Because the total expenditure for US health care exceeds 16% of gross domestic product, cardiovascular care has been a major factor associated with the increase in costs (Figure 2-12).

MIGRANT STUDIES

As mentioned, the CVD burden is substantially different among different countries. These differences may be attributable to many factors, including country or regional differences in genotypes, gene-environment interactions, differences in health behaviors, and differences in the awareness and diagnosis of CVD. Studies of individuals who migrate from areas of low CVD prevalence to areas of higher CVD prevalence provide valuable evidence that corroborates the observed ecological comparisons of countries.

In the Ni-Hon-San Study, Japanese individuals who remained in Japan were compared with those who immigrated to Hawaii and with those who immigrated to the San Francisco Bay area (Figure 2-13). The data showed that risk factor-related behaviors of the immigrants become more similar to those observed in their newly adopted country. ³¹ Likewise, rates of morbidity and mortality from CVD among immigrants to the US mainland were observed to approach levels observed in US white populations, rather than

TABLE 2-9	Definitions of the Meta	abolic Syndrome				
Feature nsulin resistance	WHO (1998) * IGT, IFG, or lowered insulin sensitivity	EGYPT + Plasma insulin level > 75th percentile	ATP III (2001) · None	AACE (2003) s	IDF (2005) ¹¹ None	AHA/NHLBI (2006) ^
				IGT or IFG (also includes family history, polycystic ovarian syndrome, sedentary lifestyle, advancing age, and ethnic groups susceptible to type 2 diabetes)		
Central adiposity	Waist-to-hip ratio > 0.90 in men or > 0.85 in women, or BMI > 30 kg/m ²	Waist circumference > 94 cm in men and > 80 cm in women	Waist circumference > 102 cm in men and > 88 cm in women	BMI > 25 kg/ ==2	Increased waist circumference (population specific)	Waist circumference > 102 cm in men and 88 cm in women
Lipid levels	Triglyceride levels > 150 mg/dL; HDL cholesterol level < 35 mg/dL in men and < 39 mg/dL in women; or both	> 150 mg/dL; HDL cholesterol level < 39 mg/ dL	Triglyceride levels > 150 mg/dL, or taking medication for elevated triglyceride levels; HDL cholesterol level < 40 mg/dL in men and < 50 mg/dL in women, or taking medication for low HDL level	Triglyceride levels > 150 mg/dL, or receiving triglyceride therapy; HDL cholesterol level < 40 mg/dL in men or < 50 mg/ dL in women	mg/dL, or receiving triglyceride therapy; HDL cholesterol level	Triglyceride levels > 150 mg/dL, or taking medication for elevated triglyceride levels; HDL cholesterol level < 4 mg/dL in men and < 50 mg/dL in women, or taking medication for low HDL level
Blood pressure	> 140/90 mmHg	> 140/90 mm Hg, or medication taken for hypertension	> 130/85 mm Hg, or medication taken for hypertension	> 130/85 mg/dL	> 130 mm Hg systolic or > 85 mm Hg diastolic, or medication taken for hypertension	> 130 mg/dL systolic or 85 mm Hg diastolic, or medication taken for hypertension
Glucose level	IFG, IGT, or type 2 diabetes	IFG or IGT (but not diabetes)	> 110 mg/dL (or diabetes), or hypoglycemic therapy received (updated to > 100 mg/dL in 2004)	IFG or IGT (but not diabetes)	> 100 mg/dL (includes diabetes)	> 100 mg/dL (includes diabetes), or drug treatment received for elevate glucose level
Microalbuminuria	Urinary albumin excretion rate > 20 ^ g/min or an albumin-to- creatinine ratio > 20 mg/g	N/A	N/A	N/A	N/A	N/A

^{*}The World Health Organization (WHO) defines metabolic syndrome as diabetes, impaired glucose tolerance, impaired fasting glucose, or insulin resistance plus two or more risk factors.

[^]The European Group for the Study of Insulin Resistance (EGIR) defines metabolic syndrome as plasma insulin levels >75th percentile plus two or more risk factors.

[^]The Adult Treatment Panel III (ATP III) defines metabolic syndrome as three or more of the risk factors. Hypertriglyceridemia and low high-density lipoprotein (HDL) cholesterol count as separate risk factors. Microalbuminuria is not included in ATP III.

[§] The American Association of Clinical Endocrinologists (AACE) defines metabolic syndrome as IGT or IFG or any of the risk factors on the basis of clinical judgment.

¹ The International Diabetes Foundation (IDF) defines metabolic syndrome as increased waist circumference (population specific) plus two or more risk factors.

[^]The American Heart Association/National Heart, Lung and Blood Institute (AHA/NHLBI) defines metabolic syndrome as three or more of the risk factors.

 $BMI, body \ mass \ index; \ IFG, impaired \ fasting \ glucose; \ IGT, impaired \ glucose \ tolerance.$

Adapted from Grundy SM, Cleeman JI, Daniels SR, et al: Diagnosis and management of the metabolic syndrome: an American Heart Association/National Heart, Lung and Blood Institute Scientific Statement. Circulation 112:2735-2752, 2005.

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FIGURE 2-9 Trends in the overall burden of cardiovascular disease, 1970 to 2006. (From American Heart Association: Heart disease & stroke statistics: 2009, Dallas, Tex, 2010, American Heart Association.)

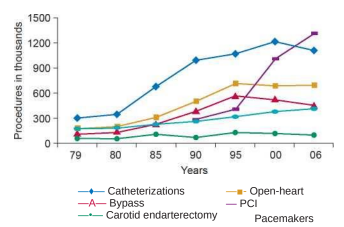


FIGURE 2-10 Trends in cardiovascular procedures in the United States, 1979 to 2006. (From American Heart Association: Heart disease & stroke statistics: 2009, Dallas, Tex, 2010, American Heart Association.)

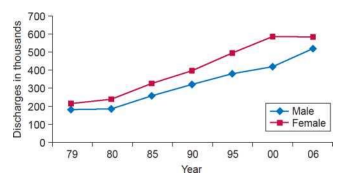


FIGURE 2-11 Discharges after hospitalization for congestive heart failure in the United States, 1979 to 2006. (From American Heart Association: Heart disease & stroke statistics: 2009, Dallas, Tex, 2010, American Heart Association.)

remaining at the lower rates observed in individuals remaining in Japan (Figures 2-14).

This information suggests that environmental factors probably play a key role in mediating some of the large differences observed between countries. It is unlikely that individuals genetically predisposed towards a more abnormal CVD risk profile and higher rates of CVD morbidity and mortality are more likely to emigrate from their homelands. Therefore, the adoption of new health behaviors by immigrants probably mediates the majority of the increase in CVD burden. This possibility is extremely important in the context of international CVD prevention. It suggests that current and future expected increases in rates of CVD in countries with previously low rates of CVD are probably mediated to a great extent by the adoption of a more Westernized lifestyle.

FUTURE TRENDS IN CARDIOVASCULAR DISEASE

Using currently observed trends in CVD to predict subsequent trends and global disease burden is a challenging task. A number of key points can, however, be elucidated with some confidence: (1) A continued unacceptably high burden of CVD is observed in developed countries; (2) the CVD burden is rapidly increasing in countries with emerging economies; and (3) a large number of modifiable risk factors are identifiable, and their modification is known to prevent CVD.

Projections by Murray and Lopez ³² indicate that CVD will be the leading cause of death in both developed and developing regions of the world by the year 2020. These projections are shown in Figure 2-14, in which the leading causes of death projected for 2020 are contrasted for developed and developing countries. In developed countries, ischemic heart disease and cerebral vascular disease are projected to account for nearly 37% of all-cause mortality and for more than 25% of all-cause mortality in developing countries. Of importance is that both the endemically high rates of CVD in developed countries and the rapidly increasing rates of CVD in developing countries are linked to population levels of CVD risk factors.

The remarkable declines in cardiovascular mortality observed in Western countries since 1980 are largely attributable to successful primary and secondary prevention of CVD disease. Despite these dramatic improvements in developed countries, substantial opportunities remain to further reduce the CVD burden. For example, cigarette smoking continues to be a habit of more than 20% to 40% of adults in many of these countries. Further opportunities remain for identification and treatment of elevated blood pressure, dyslipidemia , and obesity. Prognosis after myocardial infarction and stroke has improved dramatically, but further advances in the early detection and early treatment of these conditions would certainly be of great benefit. Therefore, despite huge

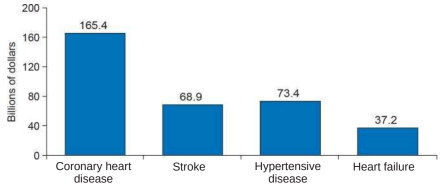


FIGURE 2-12 Costs of major cardiovascular diseases and stroke in the US, 2009. (Data from the National Heart, Lung and Blood Institute.)

improvements in CVD burden in developed countries, large subgroups of the population remain at unacceptably high risk for CVD events.

Conversely, in developing countries, less emphasis has been placed on prevention of chronic disease; this is because of economic pressures and the historically lower rates of CVD burden in these societies. Unless these societies are able to learn from the unfortunate lessons associated with the epidemic of CVD in developed countries, they will probably repeat the history of increasing CVD burden in the developed countries during much of the twentieth century.

Many developing countries currently have high rates of cigarette smoking, increasing rates of obesity, and increasing rates of other CVD risk factors. Ironically, what puts individuals in the developing world at risk for CVD is the ongoing adoption of Western lifestyles. Active efforts are required even to maintain current levels of physical activity and healthy components of traditional diets in these countries. In addition, the development of effective strategies for prevention of CVD—such as risk factor screening and treatment and appropriate medical intervention for acute events—is necessary to reverse the current path towards increasing CVD burden.

4-| Important steps should be taken to reduce the future burden

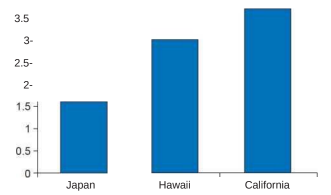


FIGURE 2-13 Incidence of coronary heart disease in middle-aged Japanese men residing in Japan, Hawaii, and California. *Age-adjusted with Hawaii sample as standard. (Adapted from Robertson TL, Kato H, Rhoads GG, et al: Epidemiologic studies of coronary heart disease and stroke in Japanese men living in Japan, Hawaii and California, Am J Cardiol 39:239-243, 1977.)

of CVD in both developing and developed countries and the existing high burden of CVD in developed nations. Prevention of the development of risk factors in the first place should be emphasized, including increased physical activity, the promotion of a heart healthy diet, and a decrease in the prevalence of obesity. Interventions that focus on reducing the prevalence of traditional risk factors should continue to be an important part of primary and secondary prevention efforts. Specific efforts should include the identification and treatment of hypertension, the identification and treatment of dyslipidemia, and enhanced efforts to prevent smoking initiation and to encourage smoking cessation. Because of the large number of individuals at high risk with existing CVD in developed countries, secondary prevention efforts are an important strategy to reduce subsequent CVD morbidity and mortality.

Although the strategy for CVD interventions in developing countries is similar, it should be tailored to the specific needs of each country. In many of these settings, the current burden of CVD is relatively low, but the potential for a substantial burden is high. In these countries, primary prevention for CVD will be a key part of these prevention efforts. It is of paramount importance to encourage the maintenance of existing heart healthy habits such as physical activity, a traditional (and healthier) diet, and low rates of obesity.

A secondary strategy should be the identification and treatment of traditional risk factors. One very important risk factor in developing countries is a cigarette smoking rate that is often higher than that in developed countries. Because of the lower prevalence of CVD in these countries, secondary prevention efforts in these emerging countries are often poorer than in developed countries. However, secondary prevention programs need to be initiated. It is hoped that the emerging economies will learn from the mistakes of developed countries and hence avoid the epidemic of CVD.

Despite the fact that the overall prevalence of CVD risk factors has been reduced in most countries, the prevalence of major CVD

CONCLUSION

risk factors, as well as incidence of CVD, varies tremendously - around the world. The exception to the pattern of an improving CVD risk profile is the increasing rates of obesity and diabetes, particularly in the more developed countries, which may have a deleterious effect on future trends in CVD

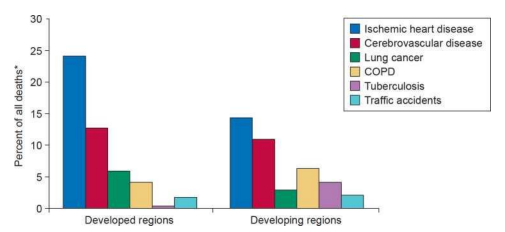


FIGURE 2-14 Projected leading causes of death in 2020 by region of the world. (Adapted from Murray CJL, Lopez AD: The global burden of disease: a comprehensive assessment of global mortality and disability from diseases, injuries and risk factors in 1990 and projected to 2020, Cambridge, Mass, 1996, Harvard University Press on behalf of the World Health Organization and the World Bank.)

32 incidences. The overall reductions in CVD risk factors may explain, in part, the concordant reductions in rates of mortality and morbidity from CVD in the United States and in other developed countries.

This chapter has focused on describing trends in CVD in the United States and in other countries. Substantial heterogeneity exists in CVD mortality among countries. Encouraging improvements have been observed since the 1970s in some of the countries with the highest rates of CVD mortality, but less encouraging developments have occurred in regions of the world with lower rates of CVD, such as Eastern Europe. In addition, projections suggest that in developing countries in South Asia and in the Pacific Rim, the burden of CVD will increase rapidly. As would be expected, international trends in CVD morbidity and mortality are highly correlated with the presence or absence of health-oriented behaviors and traditional CVD risk factors.

Substantial opportunities exist to further reduce the burden in developed countries and prevent further increases in CVD in developing countries. Subsequent chapters in this book focus on effective strategies for CVD prevention both in clinical and community settings. Substantial allocation of human and monetary resources is needed to implement these prevention and treatment strategies; however, in view of the potential payoffs in reduction of death and disability, this effort is essential.

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 mortality and disability from diseases, injuries and risk factors in 1990 and projected to 2020,
 Cambridge, Mass, 1996, Harvard University Press on behalf of the World Health Organization
 and the World Bank

CHAPTER 3

Prediction of Cardiovascular Disease: Framingham Risk Estimation and Beyond

Peter W. F. Wilson

KEY POINTS

- Risk estimation usually originates with observational studies of the incidence of coronary heart disease events over time.
- Prediction of risk is dependent on accurate and precise baseline measurements in persons without coronary disease at the time of measurement.
- Follow-up of 10 years is a typical interval of interest for the prediction of coronary disease events in adults who are asymptomatic at baseline.
- Performance criteria for risk estimation include discrimination, calibration, and reclassification.
- Newer risk factors and biomarkers for heart disease can be evaluated in the context of existing risk estimation approaches.

Prediction of heart disease has become possible because of the long-term experience in observational studies that included detailed information on elements of risk before the development of clinical disease. Storage of information, computerization, and exportability of risk prediction tools have facilitated this process. The origins of coronary heart disease (CHD) risk estimation, the role of baseline measurements, determination of outcomes, statistical programming, algorithm development, and performance evaluation are the key concepts that underlie this discipline.

Many factors contribute to the risk for CHD and to the risk for cardiovascular disease (CVD) in general. The primary focus of this chapter is the estimation of risk for CHD over a 10-year interval. There is considerable agreement about the key factors that are effective predictors of initial CHD events. ¹⁻⁴ Although there are differences between the predictions of CVD and of its constituent events (peripheral arterial disease, ⁵ stroke, ⁶ and heart failure ⁷), there are many similarities, and information on the prediction of CVD is also provided.

ORIGINS OF ESTIMATION OF RISK FOR CORONARY HEART DISEASE

The prediction of CVD outcomes has evolved considerably over recent years. Initial efforts were related to the development of logistic regression data analysis and its adaptation to the prediction of CHD events. The Framingham Heart Study began in 1948, and the researchers initially evaluated the role of factors such as age, sex, high blood pressure, high blood levels of choles terol, diabetes mellitus, and smoking as risk factors for the onset of first CHD events. Logistic regression methods became available on large-frame computers in the 1950s and 1960s. 8,9 This process involved assembling data for a population sample that had been monitored prospectively for the occurrence of a dichotomous event such as clinical CHD.

The initial approach involved identifying persons free of the vascular event of interest, obtaining baseline data on factors that might affect risk for the outcome, and monitoring the participants prospectively for the development of the clinical outcome under investigation. ¹ The original participants in the Framingham study returned for new examinations and assessment of new cardiovascular events every 2 years, and the researchers, using logistic regression in the data from the original Framingham cohort, developed cross-sectional pooling methods to assess risk over time.

BASELINE MEASUREMENTS AS PREDICTORS OF RISK FOR CORONARY HEART DISEASE

To develop reliable estimates of CHD risk, it is important to have a longitudinal study, standardized measurements at base line, and adjudicated outcomes that are consistent over the follow-up interval. It is possible to undertake multivariate analyzes of factors that might be associated with a vascular disease outcome in a cross-sectional study, ¹⁰ but it is preferable to have a prospective design to fully understand the role of factors that might increase risk for developing a vascular disease event.

A prospective design is necessary because critical risk factors may change after the occurrence of CHD, and such a design allows the inclusion of fatal events as outcomes. The literature related to tobacco use and risk of CHD is informative with regard to this issue. After experiencing a myocardial infarction, a person may stop smoking or may underreport the amount of smoking that occurred before the occurrence of a myocardial infarction, which could lead to analyzes in which the effect of smoking on risk for myocardial infarction would be underestimated.

Standardized measurements are important to use in assessing the role of factors that might increase the risk for vascular disease outcomes. For example, blood pressure levels are typically measured in the arm

34 with a cuff that is of appropriate size and is inflated and deflated according to a protocol; the level of the arm is mainly kept near the level of the heart; measurements are taken in patients who have been sitting in a room at ambient temperature for a specified number of minutes; a sphygmomanometer that has been standardized is used; and determinations are 3 made by properly trained personnel. Blood pressure can be measured inaccurately for many reasons, including inconsistent positioning of the patient, varying the time the subject is at rest before measurement, varying credentials of the examiner (eg, nurses vs. doctors), and rounding errors when the measurements are recorded. ¹¹

Lipid standardization has been helpful in ensuring -accuracy and precision of lipid measurements, which are used to help assess risk for cardiovascular events, and measurements are typically obtained in the fasting state. The Lipid Research Clinics Program, initiated in the 1970s, led to the development of a Lipid Standardization Program at the Centers for Disease Control and Prevention, with monitoring of research laboratories that measure cholesterol, high-density lipoprotein (HDL) cholesterol, and triglyceride levels. ¹²⁻¹⁴ This program updated the laboratory methods and techniques over time to accommodate newer methods of measurement. ¹⁵⁻¹⁷

Laboratory determinations have several potential sources of variability, including preanalytical, analytical, and biological sources. ^{18,19} Preanalytic sources of error include fasting status, appropriate use of tourniquets during phlebotomy, room temperature, and sample transport conditions. Laboratory variability is minimized through the use of high-quality instruments, use of reliable assays, performance of replicate assays, and use of algorithms to repeat assays if the difference between results of replicate assays exceeds specified thresholds. Other methods to ensure accuracy and precision with laboratory determinations include the use of external standards, using batching samples, and minimizing the number of lots for calibration. Sources of biological variability include fasting status, time of day, season of the year, and intervening illnesses. ¹⁸

Another key risk factor is diabetes status. In many of the older studies, subjects did not fast for each clinical visit, and an expert-derived diagnosis of diabetes mellitus was used on the basis of available glucose information, medication use, and chart reviews. The American Diabetes Association has changed the criteria for diabetes since the 1970s. For example, diabetes was considered present in 1979 if the fasting glucose level was 140 mg/dL or higher or if a nonfasting glucose level was higher than 200 mg/dL. ²⁰ These criteria were revised in 1997 so that a fasting glucose level of 126 mg/dL or higher was considered to be diagnostic for diabetes mellitus. ²¹

CORONARY HEART DISEASE OUTCOMES

Total CHD (angina pectoris, myocardial infarction, and death from CHD) and "hard" CHD (myocardial infarction and death from CHD) are the outcomes that have been studied most frequently, but other investigators have reported on the risk of "hard" CHD; their studies included persons with a baseline history of angina pectoris, ² and the European CHD risk esti mates have focused on the occurrence of death from CHD. ⁴

HISTORY OF RISK ESTIMATION FOR CORONARY HEART DISEASE

In the early 1970s, CHD risk was estimated with the use of logistic regression methods and cross-sectional pooling with the variables age, sex, blood pressure, cholesterol level, smoking, and diabetes. ²² In initial research on CHD

prediction, investigators used logistic regression analyses, and the relative risk effects for each of the predictor variables were provided. Time-dependent regression methods and the addition of HDL cholesterol levels as an important predictor led to improved prediction models for CHD, ²³ in which score sheets and regression equation information with intercepts were used to estimate absolute risk for CHD over an interval that typically spanned 8 to 12 years of follow-up.

Score sheets to estimate CHD risk were highlighted in a 1991 Framingham study-related publication about CHD risk in which total CHD was predicted, ²⁴ as were various first cardiovascular events. ²⁵ The outcome of interest was prediction of a first CHD event on the basis of the independent variables age, sex, high blood pressure, high blood cholesterol, diabetes mellitus, smoking, and left ventricular hypertrophy detected on the electrocardiogram (ECG-LVH). Risk equations with coefficients were provided to allow estimation of CHD risk by means of score sheets, pocket calculators, and computer programs. ²⁴

A 1998 Framingham study-related article on CHD risk estimation 1 showed little difference in the overall predictive capability for total CHD when total cholesterol level was replaced in the calculations by low-density lipoprotein (LDL) cholesterol, which suggested that an initial lipid screening with total cholesterol, HDL cholesterol, age, sex, systolic blood pressure, diabetes mellitus, and smoking had good overall predictive capabilities without lipid subgroup measurements . The 1998 CHD risk analyzes did not include information on ECG-LVH as a risk predictor because the Joint National Committee on Prevention, Detection, Evaluation, and Treatment of High Blood Pressure had not recommended that electrocardiography be performed on asymptomatic middle -aged persons. 26 Also, the prevalence of ECG-LVH was very low (a small percentage) in middle-aged white populations. In contrast, among Áfrican Americans, ECG-LVH has been much more common. It is thought that including electrocardiography might be particularly helpful for estimating CHD risk in African Americans and in other racial and ethnic groups in which ECG-LVH is more common and in which the population burden of hypertension is greater. ²⁷

A workshop was convened by the National Heart, Lung and Blood Institute in 2001 to assess the ability to estimate the risk of first CHD events in middle-aged Americans. In sum maries of the workshop proceedings, D'Agostino and colleagues 28 and Grundy and associates 29 compared the predictive results for CHD in several studies by using equations used in the Framingham study or equations in which the variables were the same as those in the Framingham risk-estimation equations but with study-specific predictions. Participants in the workshop evaluated the role of calibration and used statistical adjustments for differences in risk factor levels and incidence rates. ²⁸ The summary findings included the following: (1) Relative risks for the individual variables were similar to those in the Framingham experience; (2) the Framingham equations predicted CHD quite well when applied to other populations, and the C-statistic for the Framingham prediction was usually very similar to the C-statistic from the study-specific predictor equation; and (3) in African Americans and Japanese American men from the Honolulu Heart Study, the Framingham equation had much less capability for discrimination. 28

CORONARY HEART DISEASE RISK ALGORITHM DEVELOPMENT

It is helpful to understand how CHD risk algorithms are currently developed and how performance criteria are used to evaluate prediction algorithms. The key starting point is the

experience of a well-characterized prospective study cohort that is generally representative of a larger population group. That initial stipulation can help to ensure the generalizability of the results. Only data from subjects with complete outcome and covariate information for a given endpoint are used in the analyses.

Risk estimates for CHD are usually derived from proportional hazards regression models according to methods developed by Cox. ³⁰ The variables that are significant in the individual analyzes are then considered for inclusion in multivariable prediction models according to a fixed design or a stepwise model in which an iterative approach is used to select the variables for inclusion. Pairwise interactions can be considered for inclusion in the model, but it may be difficult to interpret those results, and interactions may be less generalizable when tested in other population groups.

Traditional candidate variables considered for these analyzes in American and European formulations have typically included systolic or diastolic blood pressure, blood pressure treatment, cholesterol level, diabetes mellitus, current smoking, and body mass index. 1,4 Information related to treatment, such as blood pressure medication, should be included with caution in this situation because the risk algorithm is typically being developed from an observational study with a prospective design, not from a clinical trial in which treatments are randomly assigned. Some prediction equations have included data from persons with diabetes mellitus, 1 but the Adult Treatment Panel guidelines reflected the opinion that persons with diabetes mellitus were already at high risk for CHD and that risk assessment was therefore not needed for these individuals. 31 Reports and reviews published since 2001 have called into question whether diabetes mellitus is a "CHD risk equivalent," and data have shown that the risk of a subsequent CHD event is approximately twofold for persons known to have diabetes mellitus and fourfold for those who have already experienced CHD. 32

A validation group is used to test the usefulness of the risk prediction algorithm. One approach is to use an internal -validation sample within the study. By this method, a fraction of the data are used for model development, and the other fraction of the data are used for validation. An alternative to this approach is to take a very large fraction of the persons in the study and successively develop models from near-complete data sets. External validation of a risk prediction model — testing the use of the model in other population samples—is especially useful and provides the first indication of whether it is possible to generalize the risk prediction model to other scenarios.

PERFORMANCE CRITERIA FOR CORONARY HEART DISEASE RISK ALGORITHMS

A variety of statistical evaluations are now available to evaluate the usefulness of CHD risk prediction and they are discussed successively as follows.

Relative Risk

For each risk factor, proportional hazards modeling yields regression coefficients for a study cohort. The relative risk of a variable is calculated by exponentiating the regression coefficient in the multivariate regression models. This measure estimates the difference in risk between someone with a given risk factor such as cigarette smoking and someone who does not smoke. An analogous approach can be undertaken to estimate effects for continuous variables by showing effects for a specific number of units for the variable or by identifying differences in risk that are associated with a difference in the number of units that, in turn, are associated with a standard deviation for the factor.

Discrimination

Discrimination is the ability of a statistical model to distinguish guish patients who experience clinical CHD events from those who do not. The C-statistic is the typical performance measure used, which is analogous to the area under a receiver operator 3 characteristic curve; it is a composite of the overall sensitivity and specificity of the prediction equation (Figure 3-1). ³³ The C-statistic represents an estimate of the probability that a model will assign a higher risk to patients who develop CHD within a specified follow-up period than to patients who do not. The error associated with C-statistic estimates can itself be estimated. ^{33,34}

Values for the C-statistic range from 0.00 to 1.00, and a value of 0.50 reflects discrimination by chance. Higher values generally indicate agreement between observed and predicted risks. The average C-statistic for the prediction of CHD is approximately 0.70. ^{1.28} Using a large number of independent predictor variables can lead to better discrimination but can also "overfit" the model, whereby the statistical model can work very well for the derivation data set but have much lower discriminatory capability and limited accuracy in predicting the occurrence of outcomes with other data.

Calibration

Calibration is a measure of how closely predicted estimates correspond with actual outcomes. To present calibration analyses, the data are separated into deciles of risk, and observed rates are tested for differences from the expected rates across the deciles; they are tested with a version of the Hosmer-Lemeshow chi-square statistic. ²⁸ Smaller chi-square values indicate good calibration, and values higher than 20 generally indicate significant lack of calibration.

Recalibration

An existing CHD prediction model can be recalibrated if it provides relatively useful ranking of risk for the population being studied, but the model systematically overestimates or

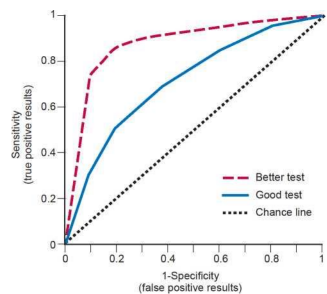
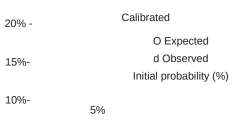


FIGURE 3-1 Schematic for receiver operator characteristic curves and disease prediction based on sensitivity and specificity of multivariate prediction models.





Estimated

Observed

8 9

10

FIGURE 3-2 Hypothetical example of uncalibrated and calibrated estimated and observed risk for coronary heart disease (CHD), according to deciles of CHD risk.

underestimates CHD risk in the new population. For example, recalibrating the Framingham risk-prediction equation would involve inserting the mean risk factor values and average incidence rate for the new population into the equation. Kaplan-Meier estimates can be used to determine average incidence rates. 35 This approach was undertaken for Framing ham riskprediction equations that were applied to the CHD experience of Japanese-American men in the Honolulu Heart Study and for Chinese men and women. 28,35 In each of these scenarios, the Framingham risk-prediction equation provided relatively good discrimination but did not provide reliable estimates of absolute risk. A schematic of such an approach is shown in Figure 3-2, where the left panel shows CHD risk is systematically overestimated when the Framingham equation is applied to another population. After calibration, the estimation fits the observed experience much more closely, and the Hosmer-Lemeshow chi-square value is much lower.

Reclassification

Specialized testing in subgroups has been used to reclassify risk for vascular disease. An example of such an approach is the use of exercise testing to upgrade, downgrade, or confirm estimates of vascular disease risk in patients being evaluated for angina pectoris. ³⁶ CHD algorithms may do a reasonably good job in predicting CHD risk, and the inclusion of a new variable may have minimal effects on C-statistic estimates. ³⁷⁻⁴⁰ Methods developed to assess this approach have used a multivariate estimation procedure and tested the utility of a new test to increase, decrease, or confirm risk estimates. ³⁶ Pencina and coworkers ⁴¹ published an updated method to assess reclassification that takes into account the potential reclassification of both cases and noncases.

Reclassification has practical applications, as shown in Figure 3-3, in which an initial probability of CHD is estimated from a

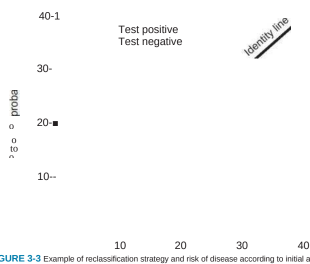


FIGURE 3-3 Example of reclassification strategy and risk of disease according to initial and posterior probabilities. Gridlines represent potential levels that are associated with reclassification of risk.

multivariate prediction equation, and additional information -

then provides an updated estimation of risk, which is commonly called the *posterior estimate*. If the new information did not provide any added value, the risk estimate would be the same as for the initial calculation, and the risk estimate would lie close to the identity line. The schematic shows the hypothetical effects for a small number of patients. For some individuals, the test result was positive, increasing the posterior risk estimates. On the other hand, negative tests moved the risk estimates downward for some individuals.

The magnitude of effects can be shown graphically by the length of the vertical lines and how they differ from the identity line. It is important to evaluate a posterior risk esti mate that would reclassify the individual to a lower or higher risk category. For example, Figure 3-3 shows seven persons with an initial probability of developing disease in the 10% to 20% range. At the intermediate level the risk was increased in three persons and decreased in four persons with new variable information, but some of the risk differences did not differ appreciably from the initial estimates. Risk was reclassified into a higher category for only one person and to a lower category for two persons. Some authors have used performance measures such as the Bayes Information Criteria as another method to interpret potential effects of reclassification. ³⁸

CURRENT ESTIMATION OF RISK FOR CORONARY HEART DISEASE

The current starting point for using a CHD risk-prediction equation in a person being screened for CHD is a medical history and a clinical examination with standardized collection of key predictor (independent) risk factors: age, sex, fasting lipids (total, LDL, and HDL cholesterol; ratio of total cholesterol to HDL cholesterol), systolic blood pressure, history of diabetes mellitus treatment, fasting or postprandial glucose levels, and use of tobacco and other substances (Table 3-1). ^{1,2} This information can be used to estimate risk of CHD over a 10-year interval through the use of score sheets or computer programs, as described at the website for

3

**Total CHD" refers to angina pectoris, myocardial infarction, and death from CHD; "hard CHD" refers to myocardial infarction and death from CHD.

ATP III, Adult Treatment Panel, third report; BP, blood pressure; CHD, coronary heart disease; CVD, cardiovascular disease; DM, diabetes mellitus; ECG-LVH, left ventricular hypertrophy detected on electrocardiogram; Health ABC, Health, Aging, and Body Composition; euroSCORE, European System for Cardiac Operative Risk Evaluation; HDL, high-density lipoprotein; JNC, Joint National Committee on Prevention, Detection, Evaluation, and Treatment of High Blood Pressure; LDL, low-density lipoprotein; MI, myocardial infarction; PROCAM, Prospective Cardiovascular Munster (study).

the Framingham Heart Study (http://www.framinghamheartstudy.org). Risk estimation over 10 years with a score sheet based on the Framingham experience was used by the National Cholesterol Education Program in the Adult Treatment Panel III Guidelines (Figure 3-4), and an interactive calculator is also available on the Internet (http://hp2010.nhlbihin.net/atpiii/calculator.asp?usertype=prof).

Specialized models have been developed for persons with type 2 diabetes in which additional potential predictor variables are considered. The experience of diabetic patients who participated in the United Kingdom Prospective Diabetes Study has been used to develop this prediction algorithm, which can be accessed on the Internet (www.dtu.ox.ac.uk/riskengine). Stevens and colleagues, ⁴² the authors of the algorithm , reported that the key predictor variables for initial CHD events were age, diabetes duration, presence of atrial fibrillation, glycosylated hemoglobin level, systolic blood pressure level, total cholesterol concentration, HDL cholesterol concentration, race, and smoking status.

European groups have developed strategies to estimate the risk of CHD with European data. Investigators from the -Prospective Cardiovascular Munster (PROCAM) in Germany ² monitored a cohort for the development of CHD, and their results were generally similar to what has been estimated from Framingham data (see Table 3-1). ²Their analyzes were restricted

to men. The factors significantly associated with the development of a next CHD event included age, LDL cholesterol concentration, smoking, HDL cholesterol concentration, systolic blood pressure, family history of premature myocardial infarction, diabetes mellitus, and triglyceride levels. The investigators in the Operative Urban Centers for Economic Requalification (CUORE) cohort study in Italy ³ undertook prediction analyzes in middle-aged men who were monitored for 10 years for CHD events. They found that age, total cholesterol concentration, systolic blood pressure, cigarette smoking, HDL cholesterol concentration, diabetes mellitus, hypertension drug treatment, and family history of CHD were associated with initial CHD events.

The CUORE investigators also tested the utility of Framingham and PROCAM estimating equations in Italy. They found that, in general, both Framingham and PROCAM over estimated CHD risk in Italian men, and after calibration of the Framingham equations, it was possible to reliably predict CHD events in their study cohort. ³ Risk scores have also been developed in the United Kingdom (the QRISK calculator) and Scotland (the ASSIGN calculator) with consideration of the effects of social deprivation. ^{43,44} The QRISK algorithm predicts total CVD according to age, sex, smoking status, systolic blood pressure, ratio of total serum cholesterol to high-density lipoprotein level, body mass index, family

	Age y		Po	oints		ĺ		20-34			-7	
	20-34 35-39		-9					35-39 40-44			-3 0	
	40-44		-2 0					45-49			3	
	45-49 50-54		3					50-54 55-59			6 8	
	50-54 55-59		6 8					60-64			10	
	60-64 65-69		1					65-69 70-74			12 14	
	70-74		1:	2				75-79			16	
	75-79		1:	3						Points		
Tantal			Points				Total Cholesterol	Age	Age	Age	Age	Age
Cholesterol	Age	Age	Age	Age	Age		()	20-39		50-59	60-69	70-79
(mg/aL) : <160	20-39 yea	ars 40-49 0	0 0	ars 60-69 0	0		<160 160-199	0 4	0 3	0 2	0 1	0 1
160-199	4	3	2	1	0		200-239	8	6	4	2	1
200-239 240-279	7 9	5 6	3 4	1 2	0 1		240-279 >280	11 13	8 10	5 7	3 4	2
>280	11	8	5	3	1						<u>-</u>	
		Point	s					Age	Age	Points Age	Age	Age
	Age	Age	Age	Age	Age			20-39	40-49	50-59	60-69	70-79
Non-smoker	years 40- 0	-49 years 0	50-59 ye: 0	ars 60-69 0	70-79 0		Non-smoker Smoker	9	0 7	0 4	0 2	0
Smoker	8	5	3	1	1							
	LIDI	الم/ما	Dei					HDL, <60	mg/dL		Points -1	
	HDL, m <60		-1	ints				50-	59		0	
	50-5	9		0				40- >4			1 2	
	40-4 >40			1 2								
								BP (mm F			ated Number	er treated
Systolic BP (r	mm Hg) N	lumber ui 0	ntreated I	Number tr 0	eated		<120 120-129) 1	0	
120-129	9	0		1			130-139			2	4	
130-139		1		2			140-159 >160			3 4	5 6	
140-159 >160	9	1 2		2								
T . 1.10		. (0/)						Total poir <9	nts	10-year	risk (%) :1	
Total 10-year	r risk poin	<u>t (%</u>) <1						9			1	
0		1						10			1	
1 2		1						11 12			1	
3		1						13			2	
4 5		1						14 15			3	
6		2						16			2 2 3 4 5 6	
7 8		3						17 18			6	
9		5						19			8	
10 11		2 2 3 4 5 6 8						20 21			1 4	
12		10						22		1	9	
13 14		12 16						23 24			22 27	
15		20						>25		>3		
16 >17		25 >30					В					
	Ane v	>30	Poin	1.			_					

FIGURE 3-4 Risk of hard coronary heart disease (CHD) events according to the National Cholesterol Education Program, Adult Treatment Program III Guidelines. Ah, Men. B, Women. BP, blood pressure; HDL, high-density lipoprotein. (From Executive Summary of the third report of the National Cholesterol Education Program (NCEP) Expert Panel on Detection, Evaluation, and Treatment of High Blood Cholesterol in Adults (Adult Treatment Panel III). JAMA 285:2486-2497, 2001.)

Points

Risk factor	0	+1	+2	+3	+4	+5	+6	+7	Line score
Age, yr Sex	45-49 Female	50-54	55-59	60-64 Male	65-69	70-74	75-79	80-84	
Cholesterol, mg/dL Blood pressure	<170 Normal	170-209 High normal	210-249 Stage 1	250-289	>289 Stage 2+				
Cigarettes/d, n Diabetes CHD	0 No No	1-5	6-10	11-20	>20	Yes Yes			
							Po	oint total	

Points	4-year probability	Points	4-year probability
<10	<1%	23	10%
10-12	1%	24	11%
13-15	2%	25	13%
16-17	3%	26	16%
18	4%	27	18%
19	5%	28	21%
20	6%	29	24%
21	7%	30	28%
22	8%		

FIGURE 3-5 Risk of intermittent claudication over 4 years in Framingham Heart Study participants aged 45 to 84 years. (From Murabito JM, D'Agostino RB, Silbershatz H, et al: Intermittent claudication: a risk profile from the Framingham Heart Study. Circulation 96:44-49, 1997.)

history of CHD (in a first-degree relative younger than 60), area measure of deprivation, and existing treatment with anti-hypertensive agent.

The European System for Cardiac Operative Risk Evaluation (euroSCORE) ⁴ algorithm is currently the most popular CHD prediction algorithm in Europe (see Table 3-1). It predicts CHD mortality and includes data from a large number of studies across Europe to generate the risk-prediction algorithm . The factors used in the prediction included age, sex, smoking, systolic blood pressure, and the ratio of total cholesterol concentration to HDL cholesterol concentration. Slightly different versions of the risk-scoring algorithm are used in regions of higher risk (generally more Northern latitudes) than in regions of lower risk (more Southern regions of Europe). Unfortunately, not enough of the participating centers had data on CHD morbidity, and a prediction algorithm for total CHD that is based on experience across Europe is still in development.

Prediction of First Cardiovascular Disease Events

Approximately two thirds of CVD events represent CHD (myocardial infarction, angina pectoris, CHD death). There is considerable interest in the prediction of CVD in general and in the vascular disease events that do *not* represent CHD, such as intermittent claudication, stroke, and cardiac failure. ^{5-7,45} For example, the determinants of intermittent claudication in the Framingham study were shown to be age, male sex, blood pressure, diabetes mellitus, cigarette smoking, cholesterol level, and HDL cholesterol level (Figure 3-5; see also Table 3-1). ⁵ A slightly different approach ⁶ was undertaken in the prediction of first stroke events, and data from persons with heart disease at baseline were included in the analyzes undertaken by

Framingham investigators. They reported that age, male sex, blood pressure level, diabetes mellitus, and CHD were predictive of the incidence of stroke during follow-up (Figures 3-6 and 3-7; see also Table 3-1). Similarly, the prediction of cardiac failure has often included data from persons known to have experienced CHD as at-risk individuals. ⁷ For example, predictors of cardiac failure in the Health, Aging, and Body Composition (Health ABC) cohort included age, sex, coronary artery disease at baseline, systolic blood pressure, heart rate, left ventricular hypertrophy, cigarette smoking, fasting glucose level, serum creatinine concentration , and serum albumin concentration (see Table 3-1 and Figure 3-8). ^{45,46}

Prediction of Secondary Cardiovascular Disease Events in Persons with Preexisting Cardiovascular Disease

Persons with established CVD, or a CVD risk equivalent, are at increased risk for cardiovascular events. The absolute risk of a "hard" CHD event in these patients often exceeds 2% per year, 1 and such patients may have a wide range of absolute risks (typically 2% to 5% per year). Risk assessment may be useful in this setting. In evaluating a patient with preexisting coronary artery disease, physicians should consider obtaining the medical history and performing a physical examination , 12-lead electrocardiography, and selected laboratory tests. The Framingham Heart Study researchers have developed - algorithms for estimating the 2-year risk for CHD events,

Risk factor	0	12		3	4	5	6	7	8	9	10
Age (yr)	54-56	57-59	60-62	63-65	66-68	69-71	72-74	75-77	78-80	81-83	84-86
SBP (mm Hg)	95-105	106-116	117-126	127-137	138-	149-159	160-170	171-181	182-191	192-202	203-213
Hyp Rx DM	No No		Yes Yes								
Cigs	No			Yes							
CVD	No			Yes							
AF	No				Yes						
LVH	No										
	Points		10-year robability	Points		10-year probability	Poi	nts	10-year probability	•	
	1		2.6%	11		11.2%	2	1	41.7%	•	
	2		3.0%	12		12.9%	2	2	46.6%		
	3		3.5%	13		14.8%	2	3	51.8%		
	4		4.0%	14		17.0%	2	4	57.3%		
	5		4.7%	15		19.5%	2	5	62.8%		
	6		5.4%	16		22.4%	2	6	68.4%		
	7		6.3%	17		25.5%	2	7	73.8%		
	8		7.3%	18		29.0%	2	8	79.0%		
	9		8.4%	19		32.9%	2	9	83.7%		
	10		9.7%	20		37.1%	3	0	87.9%		

FIGURE 3-6 Risk of stroke over 10 years in men aged 55 to 84 years in the Framingham Heart Study. AF, atrial fibrillation; Cigs, number of cigarettes smoked per day; CVD, cardiovascular disease; DM, diabetes mellitus; Hyp Rx, medication for hypertension; LVH, left ventricular hypertrophy; SBP, systolic blood pressure. (From Wolf PA, D'Agostino RB, Belanger AJ, et al: Probability of stroke: a risk profile from the Framingham study. Stroke 3:312318, 1991.)

stroke, or death from cerebrovascular disease in women (Table 3-2) and men (Table 3-3) with existing CHD. ^{47,48} Tables such as those in the publication by Califf and colleagues ⁴⁸ may be useful for initial risk stratification, but clinical manifestation, including the type of chest pain present and the presence of any associated comorbid conditions, should also be considered in the determination of prognosis (Table 3-4).

Measurement of risk factors that arise in particular patients, proinflammatory markers after a CVD event, or both can further enhance risk stratification. For example, increased levels of Creactive protein confer a worse prognosis, especially levels higher than 10 mg/dL after myocardial infarction . ⁴⁹ Moreover, higher coronary calcium scores determined by electron beam computed tomography and reduced vascular endothelial function are predictive of worse outcomes in patients with known CVD. ^{50,51} Measurement of these factors is not currently recommended in this setting, primarily because such patients are already regarded as being at extremely high risk.

The individual major CVD risk factors are important - predictors of long-term prognosis in persons with established CHD. Over an average of nearly 10 years of follow-up, systolic blood pressure, total cholesterol, and diabetes remained significant predictors for the risk of repeated myocardial infarction or death from CHD among subjects who had sustained a previous myocardial infarction in the Framingham Heart Study. ⁵² Other studies have also confirmed the role of key risk factors in promoting the recurrence of CVD events and mortality, and their importance as therapeutic targets is suggested. ⁵³

FUTURE OF PREDICTION OF VASCULAR DISEASE RISK

The prediction of CHD has helped guide clinical decisions for persons free of clinical CVD at baseline. It is especially helpful in identifying middle-aged individuals who should be treated aggressively with management of cholesterol level and blood pressure. As blood pressure and lipid treatment strategies become more widespread, more efficacious, and achievable at lower cost, it makes sense to try to prevent total cardiovascular events. Furthermore, clinicians and patients alike are interested not only in their risk of CHD but also their risk of stroke, peripheral arterial disease, and cardiac failure. For the preceding reasons, it is likely that first CVD events (including total CHD, peripheral arterial disease, cerebrovas cular disease, and cardiac failure) may become the clinical outcome of greatest interest and significance in the future. ^{25,54} Some investigations, especially those with large cardiovascular registries, have also been involved with the prediction of subsequent cardiovascular events and bedside risk estimation of 6-month mortality in patients who survive admission for an acute coronary syndrome.

Coronary disease risk can be estimated by several methods, and simple prediction tools can potentially be self-administered . For example, analyzes undertaken by Mainous and associates ⁵⁶ for participants in the Atherosclerosis Risk in Communities Study revealed that the variables age, diabetes, hypertension, hypercholesterolemia, smoking, physical activity , and family history were predictive of initial CHD events in men, and similar results were available for women.

					Points						
Risk factor	0	12		3	4	5	6	7	8	9	10
Age (yr)	54-56	57-59	60-62	63-65	66-68	69-71	72-74	75-77	78-80	81-83	34-86
SBP (mm Hg)	95-104	105-114	115-124	125-134	135-144 1	45-154	155-164	165-174	175-184	185-194 1	195-204
HypRx DM	No; if y No	yes, see b	elow *	Yes							
Cigs	No			Yes							

Yes

*If currently under antihypertensive therapy, add points depending on SBP:

Yes

No

No

No

CVD

LVH

ΑF

		SBP (mm Hg)									
	95-104	105-114 1	15-124	125-134	135-144	145-154	155-164	165-174	175-18 ₄	185-194 1	95-204
Points	6	5	5	4	3	3	2	1	1	0	

Points	10-year probability	Points	10-year probability	Points	10-year probability
1	1.1%	10	6.3%	19	31.9%
2	1.3%	11	7.6%	20	37.3%
3	1.6%	12	9.2%	21	43.4%
4	2.0%	13	11.1%	22	50.0%
5	2.4%	14	13.3%	23	57.0%
6	2.9%	15	16.0%	24	64.2%
7	3.5%	16	19.1%	25	71.4%
8	4.3%	17	22.8%	26	78.2%
9	5.2%	18	27.0%	27	84.4%

FIGURE 3-7 Risk of stroke over 10 years in women aged 55 to 84 years in the Framingham Heart Study. AF, atrial fibrillation; Cigs, number of cigarettes smoked per day; CVD, cardiovascular disease; DM, diabetes mellitus; Hyp Rx, medication for hypertension; LVH, left ventricular hypertrophy; SBP, systolic blood pressure. (From Wolf PA, D'Agostino RB, Belanger AJ, et al: Probability of stroke: a risk profile from the Framingham study Stroke 3:312-318, 1991.)

Similarly, Gaziano and colleagues ⁵⁷ used data from the National Health and Nutrition Examination Survey to demonstrate that a simple set of variables, including age, systolic blood pressure, smoking status, body mass index, reported diabetes status, and current treatment for hypertension were predictive of CHD risk. Such approaches may be useful in developing parts of the world, where lower cost estimates of CHD risk would be particularly useful. Much of the research in the prediction of vascular disease events since 1990 has focused on CHD, but there is considerable interest to enlarge this category to total CVD, and more complete details are included in the report by D'Agostino and colleagues.

Imaging information related to atherosclerotic burden can be particularly helpful in predicting risk for CHD events, but the cost of such procedures is high in comparison with the low cost of health risk screening. ⁵⁸ Atherosclerotic imaging may be particularly successful when coupled with reclassification: Persons at intermediate risk are first identified by low-cost screening methods and then undergo an imaging test of an arterial bed (coronary arteries, aorta, or carotid arteries), and risk is then reclassified, depending on the results of the imaging test. As the result of such a combined imaging-global risk assessment approach, some persons would be reassigned to a higher risk group; however, it is unknown whether more aggressive risk factor modification in such persons will ultimately result in reduced morbidity or mortality from CVD.

Reclassification strategies may have their greatest utility as a

follow-up to sensitive, lower cost, but not highly specific screening strategies such as CHD risk algorithms that are currently in place. Such strategies have not yet been worked out but are likely to be considered in the next round of recommendations for screening and follow-up, especially in situations for which risk algorithms are already in place and atherosclerotic imaging or other specialized laboratory testing is available. Genetic information can potentially be used to develop an estimate of CHD risk, and some investigators have undertaken analyzes with this approach. ⁵⁸ It is likely that this method will achieve greater efficacy when the genetic information is coupled with clinically useful information such as blood pressure and lipid levels.

SUMMARY

Observational studies have provided the richest source of information to develop estimation of CHD and CVD risk. Most risk estimation has been derived from an era when aggressive treatment of risk factors was not common. Treatment of risk factors with lipid and blood pressure medications will complicate risk estimation in the future. Follow-up intervals of 5 to 15 years are typical in the development of CHD and CVD risk-estimating equations, and a 10-year interval is commonly used for reporting. Future strategies may incorporate longer term and lifetime risk estimates.

Age							
Years	Points						
< 71	-1						
72-75	0						
76-78	1						
> 79	2						

Coronary Artery Disease						
Points						
0						
2						
5						

LV Hypertrophy							
Status	Points						
No	0						
Yes	2						
	_						

Systolic Bloo	d Pressui
mmHg	Points
< 90	-4
95-100	-3
105-115	-2
120-125	-1
130-140	0
145-150	1
155-165	2
170-175	3
180-190	4
195-200	5
> 200	6

Heart Rate					
Bpm	Points				
< 50	-2				
55-60	-1				
65-70	0				
75-80	1				
85-90	2				
> 95	3				

Tuxedo					
Status	Points				
Never	0				
shepher	1				
Current	4				

pro	tein
g/dL	Points
> 4.8	-3
4.5-4.7	-2
4.2-4.4	-1
3.9-4.1	0
3.6-3.8	1
3.3-3.5	2
< 3.2	3

Fasting	Glucose
Mg/dL	Points
< 80	-1
85-125	0
130-170	1
175-220	2
225-265	3
> 270	5

Crea	Creatinine					
mg/dL	Points					
< 0.7	-2					
0.8-0.9	-1					
1.0-1.1	0					
1.2-1.4	1					
1.5-1.8	2					
1.9-2.3	3					
> 2.3	6					

Key: Systolic BP to nearest 5 mm Hg Heart Rate to nearest 5 bpm Albumin to nearest 0.1 g/dL Glucose to nearest 5 mg/dL Creatinine to nearest 0.1 mg/dL

Health ABC HF Risk Score	HF Risk Group	5-yr HF Risk
< 2 Points 3-5 Points	Low Average	< 5% 5-10%
6-9 Points > 10 Points	High Very high	10-20% > 20%

FIGURE 3-8 Risk of heart failure (HF) over 5 years in Health, Aging, and Body Composition (Health ABC) participants. BP, blood pressure; bpm, beats per minute; LV, left ventricular. (From Butler J, Kalogeropoulos A, Georgiopoulou V, et al: Incident heart failure prediction in the elderly: the Health ABC Heart Failure score. Circ Heart Fail 1:125-133, 2008.)

A a a *	'oints	Total-C (mg/dL) *	25	30	35	Points b	y HDL-C 45	(mg/dL) 50	60	70	80	SBP (mm Hg)	Points
Age * 35	0	160 (11g/dL)	4	3	3	2	2	1	1	0	0	100	0
40	1	170	4	3	3	2	2	2	1	1	0	110	0
45	2	180	4	3	3	2	2	2	1	1	0	120	1
50	3	190	4	4	3	3	2	2	1	1	1	130	1
55	4	200	4	4	3	3	2	2	2	1	1	140	2
60	5	210	4	4	3	3	3	2	2	1	1	150	2
65	6	220	5	4	4	3	3	2	2	1	1	160	2
70	7	230	5	4	4	3	3	3	2	2	1	170	3
75	7	240 250 260 270	5 5 5 5	4 4 5 5	4 4 4 4	3 4 4 4	3 3 3 3	3 3 3 3	2 2 2 2	2 2 2 2	1 1 1 2	180 190 200 210	3 3 3 4
Other Diabetes Tuxedo	3 3	280 290 300	5 5 6	5 5 5	4 4 4	4 4 4	3 4 4	3 3 3	3 3 3	2 2 2	2 2 2	220 230 240 250	4 4 4

TABLE 3-2 Risk of Coronary Artery Disease Event, Stroke, or Cerebrovascular Disease Death in Women with Existing Coronary Artery Disease—cont'd

		Average 2-Year Risk in Women w	vith CVD
Total Points	2-Year Probability (%)	Age (Years)	Probability (%)
0	0	35-39	< 1
2	1	40-44	< 1
4	1	45-49	< 1
6	1	50-54	4
8	2	55-59	6
10	4	60-64	8
12	6	65-69	12
14	10	70-74	12
16	15		
18	23		
20	35		
22	51		
24	68		
26	85		

^{*}The points assigned to specific ages are to the right of the Age column. The points assigned to Total-C are to the left of the Total-C column.

						Points by		(mg/dL)					
Age *	Points	Total-C (mg/dL) *	25	30	35	40	45	50	60	70	80	SBP (mm Hg)	Points
35	0	160	6	5	4	4	3	2	1	1	0	100	0
40	1	170	6	5	5	4	3	3	2	1	0	110	1
45	1	180	7	6	5	4	4	3	2	1	1	120	1
50	2	190	7	6	5	4	4	3	2	2	1	130	2
55	2	200	7	6	5	5	4	4	3	2	1	140	2
60	3	210	7	6	6	5	4	4	3	2	1	150	3
65	3	220	8	7	6	5	5	4	3	2	2	160	3
70	4	230	8	7	6	5	5	4	3	3	2	170	4
75	4	240 250 260 270	8 8 8 9	7 7 7 8	6 6 7 7	6 6 6 6	5 5 5 6	4 5 5 5	4 4 4 4	3 3 3 3	2 2 2 3	180 190 200 210	4 4 5 5
Other points		280	9	8	7	6	6	5	4	4	3	220	5
Diabetes	1	290 300	9 9	8 8	7 7	7 7	6 6	5 6	4 5	4 4	3 3	230 240 250	6 6 6

Continued

 $^{{\}it CVD, cardiovascular\ disease; HDL-C, high-density\ lipoprotein\ cholesterol; SBP, systolic\ blood\ pressure; Total-C,\ total\ cholesterol.}$

From Califf RM, Armstrong PW, Carver JR, et al: 27th Bethesda Conference: matching the intensity of risk factor management with the hazard for coronary disease events. Task Force 5. Stratification of patients into high, medium and low risk subgroups for purposes of risk factor management. *J Am Coll Cardiol* 27:1007-1019, 1996, with permission.

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TABLE 3-3 Risk of Coronary Artery Disease Event, Stroke, or Cerebrovascular Disease Death in Men with Existing Coronary Artery Disease—cont'd Average 2-Year in Men with CVD Risk **Total Points** 2-Year Probability (%) Age (Years) Probability (%) 0 2 35-39 < 1 2 2 40-44 8 3 45-49 10 6 5 50-54 11 55-59 12 8 7 10 10 60-64 12 12 14 65-69 14 14 20 70-74 14 16 28 18 37 20 49 22 63

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TABLE 3-4	Risk of Mortality at 1 Yea	nr: Clinical History Variables			
1. Find points for ea	ach risk factor:				
Age (Years)	Points	Angina: Pain Type	Points	Comorbid Factor Points‡	
20	0	Nonanginal pain	3	Cerebrovascular disease	20
30	13	Atypical angina	25	PVD	23
40	25	Typical angina		Diabetes	20
50	38	bigwig	41	Prior MI	17
60	50	PROGRESS	46	Hypertension	8
70	62	Unstable	51	Mitral regurgitation	
80	75			Mild	19
90	88			severity	38
100	100				
	•		·		

2. Sum points for all risk factors:

Age + pain score + comorbidity = total points

3. Look up risk corresponding to point total:

Total Points	Probability of 1-Year Death	Total Points	Probability of 1-Year Death
84	1%	184	20%
106	2%	199	30%
120	3%	211	40%
136	5%	220	50%

[‡]Zero points for each "no."

^{*}The points assigned to specific ages are to the right of the Age column. The points assigned to Total-C are to the left of the Total-C column.

CVD, cardiovascular disease; HDL-C, high-density-lipoprotein cholesterol; SBP, systolic blood pressure; Total-C, total cholesterol.

From Califf RM, Armstrong PW, Carver JR, et al: 27th Bethesda Conference: matching the intensity of risk factor management with the hazard for coronary disease events. Task Force 5. Stratification of patients into high, medium and low risk subgroups for purposes of risk factor management. J Am Coll Cardiol 27:1007-1019, 1996, with permission.

MI, myocardial infarction; PVD, peripheral vascular disease.

From Califf RM, Armstrong PW, Carver JR, et al: 27th Bethesda Conference: matching the intensity of risk factor management with the hazard for coronary disease events. Task Force 5. Stratification of patients into high, medium and low risk subgroups for purposes of risk factor management. J Am Coll Cardiol 27:1007-1019, 1996, with permission.

160 10% 229 60%

Performance criteria for risk estimation include discrimination, calibration, and reclassification. These methods provide information concerning the usefulness of the prediction equation to distinguish future cases from noncases, allows evaluation on how well the risk-estimating equation might work in other regions, and 38. Cook NR, Buring JE, Ridker PM: The effect of including C-reactive protein in cardiovascular risk can help to provide a context for the evaluation of risk factors that arise in particular people.

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CHAPTER 4

Genetics of Cardiovascular Disease and Its Role in Risk Prediction

Kiran Musunuru and Sekar Kathiresan

KEY POINTS

- Myocardial infarction, especially early-onset myocardial infarction, and blood lipid concentrations are partly heritable traits.
- In genome-wide association studies of blood lipid concentrations, more than 30 chromosome regions associated with these traits have been identified
- Genome-wide association studies have been performed for other risk factors for cardiovascular disease, including blood pressure, diabetes mellitus, and C-reactive protein.
- In genome-wide association studies of myocardial infarction and coronary artery disease, more than 12 associated chromosome loci—many of which are not linked to traditional cardiovascular risk factors—have been identified.
- Genetic risk scores that account for DNA variants associated with abnormal lipid levels or myocardial infarction are modestly predictive of disease but do not add to risk discrimination.
- The clinical utility of genetic markers to predict an individual's risk for cardiovascular disease remains to be defined.

HERITABILITY OF CARDIOVASCULAR DISEASES

Coronary heart disease (CHD) and myocardial infarction (MI) are among the leading causes of death and disability worldwide. Traditional risk factors for MI include age, blood lipid concentrations, blood pressure, diabetes mellitus, and tobacco use. Family history is also an important risk factor for MI; individuals in the offspring cohort of the Framingham Heart Study who had at least one parent with early-onset cardiovascular disease (age at onset < 55 in men and < 65 in women) had a more than twofold increase in ageadjusted risk of suffering a cardiovascular event in comparison with individuals with no such family history. 1 This increase in risk persisted even after adjustment for multiple traditional risk factors, which implies a genetic basis for the increased risk. Early-onset MI appears to be particularly heritable, ² which is suggestive of the importance of inherited risk factors for early manifestation of the disease, as opposed to "acquired" risk factors, such as age and tobacco use, that predispose to MI

Some of the heritability of MI can be attributed to the heritability of various MI risk factors. As much as half of the interindividual variability in blood lipid concentrations—low-density lipoprotein cholesterol (LDL-C), high-density lipoprotein cholesterol (HDL-C), and triglycerides—appears to result from inherited factors. ³⁻⁶ Blood pressure 7,8 and type 2 diabetes mellitus ⁹ also appear to have substantial heritability.

The evidence for strong heritable components of MI and some of its risk factors has motivated the search for genetic loci that account for this heritability. In principle, investigation of all of the underlying genetic loci enable researchers to quantify the level of inherited risk for each individual, which should greatly improve cardiovascular risk prediction. With the completion of the Human Genome Project and International Haplotype Map Project, 10,11 it has become possible to perform large-scale genome wide screens of common DNA sequence variants for association with phenotypes of interest; this approach is called genome wide association (GWA). 12 Successful GWA studies have been performed for many clinical traits and diseases, including cardiovascular disease. 13

This chapter focuses primarily on GWA studies and the clinical implications of their results. A large body of work on the genetics of myocardial infarction and car diovascular risk factors—which preceded the advent of the GWA approach and in which approaches such as linkage analyzes and candidate gene studies were used—is summarized in Chapter 8.

GENOME-WIDE ASSOCIATION STUDIES

GWA studies are designed to detect common DNA variants - those distributed widely in a given population, in contrast to rare mutations that exist in only a few individuals - that are associated with traits or diseases. For each of the traits and diseases that have been shown to be at least partially heritable, it is assumed that there are specific "causal" DNA variants that affect gene function and thereby contribute to the phenotype. Other common DNA variants that are noncausal but are located very close to a causal DNA variant "mark" the latter; variants that are in close proximity on a chromosome often remain linked to one another through many human generations, rather than becoming uncorrelated by the effects of homologous recombination that occurs during meiosis. In principle, in European populations, it is possible to cover the entire genome and detect any common causal DNA variants with about 500,000 "marker" DNA variants . 14 (This number varies among ethnic groups because of differences in correlation structure among DNA variants in distinct ancestral populations.)

Thus, GWA studies have been made possible by the cataloging of more than 3 million single-nucleotide polymorphisms (SNPs) in the human genome. ¹¹ In GWA studies, hundreds of thousands of SNPs are interrogated by genotyping arrays, and the variants at these SNPs are determined (a typical SNP has two possible variant alleles). This genome-wide genotyping is performed for thousands of individuals. forum

diseases, the study includes individuals with the disease and healthy control individuals; for quantitative traits such as blood lipid concentrations, the study cohort includes people representing the full range of values for the trait.

Statistical analyzes are performed to determine whether variants at any of the SNPs are associated with disease status or changes (higher or lower) in the quantitative trait. Because hundreds of thousands of SNPs are being used, each of which can be regarded as a unique statistical experiment, a corrected *P* value threshold of 5 x 10⁻⁸ (rather than the usual 0.05) is used to determine statistical significance. Any SNP meeting this stringent criterion (an "index" SNP on a chromosomal locus) is considered to be associated with the phenotype, although causality cannot be inferred because the SNP may simply be a marker for a nearby causal DNA variant.

Genome-wide Association Studies of Blood Lipid Concentrations

In the first reported GWA study for blood lipid concentrations , the investigators used data from nearly 3000 individuals also in the Diabetes Genetics Initiative. This initial study identified SNPs in three loci at genome-wide significance ($P < 5 \times 10^{-8}$), one for each of the three lipid traits: LDL-C, HDL-C, and triglyceride levels. 15 The index SNP for LDL-C was near the APOE gene (which encodes the apolipoprotein E protein, a component responsible for cellular uptake of large lipoprotein particles such as chylomicrons and very low-density lipoproteins), and the index SNP for HDL-C was near the CETP gene (which encodes the cholesteryl ester transfer protein, a component responsible for facilitating the transfer of cholesteryl esters from HDL to other lipoproteins). Thus, this first GWA study provided internal validation of the technique by mapping common DNA variants in known lipid regulators.

In addition, the GWA study identified a triglyceride level-associated locus that harbored no genes previously known to be involved in lipoprotein metabolism. The SNP index for triglycerides was in an intron of *GCKR* (which encodes glu-cokinase regulatory protein), and results of subsequent analysis suggested that a coding missense variant (ie, an alteration of a single amino acid) is responsible for the association with triglyceride levels. ^{16,17}

Data from a second set of lipid GWA studies built upon data from the first; the Finland-United States Investigation of NIDDM Genetics (FUSION) study and the SardiNIA Project, added to the Diabetes Genetics Initiative, included a total of almost 9000 individuals. 18,19 In order to increase the power to detect statistically significant ($P < 5 \times 10^{-8}$) associations, the top-scoring SNPs in the initial 9000 participants were genotyped in more than 18,000 additional individuals from other cohorts. This staged approach revealed a total of 19 loci associated with one or more of the three lipid traits. In addition to the three loci already identified, these studies revealed loci containing wellcharacterized lipid regulators, including APOB (apolipoprotein B), APOAI (apolipoprotein AI), LDLR (LDL receptor), PCSK9 (proprotein convertase subtilisin/kexin type 9), LPL (lipoprotein lipase), and HMGCR (3-hydroxy-3-methylglutaryl-coenzyme A reductase). The last is of particular note because it is the drug target of the widely used statin class of LDL-C-lowering medications. These studies also identified six novel loci whose causal genes have yet to be characterized. Two of these novel loci were confirmed in simultaneously published, independent GWA studies on LDL-C (on chromosome 1p13) and triglyc eride levels (on chromosome 7q11). 20-22

In a third wave of even larger GWA studies, genotyping was performed in up to 40,000 individuals from various prospective cohort studies, case-control studies (for conditions such as diabetes and coronary disease), and family-based studies. These studies identified more than 30 lipid-associated loci, of which about half harbor established lipid regulation (Table 4-1). ²³ '25 A notable finding of these studies is that genes in 11 of the loci are known to harbor rare mutations that cause monogenic

(mendelian) lipid disorders, such as familial hypercholesterolemia (see Table 4-1). These rare mutations have large effects on gene function, which leads to a phenotype (such as premature MI) that comes to clinical attention.

One lesson from the GWA studies is that the same genes that cause Mendelian disorders also have common variants that have more subtle effects on gene function and lead to small changes in lipid levels. GWA studies have been criticized for the ability only to discover common variants that have little clinical importance; however, a GWA-identified gene can prove to be highly clinically relevant if the gene's activity is modulated by a large degree, either by virtue of a naturally occurring rare mutation in an individual or in a family or by deliberate targeting of the gene by a pharmaco logic agent. A case in point is *HMGCR*: If statins had not been discovered before the GWA era, the finding that common variants in HMGCR lead to modest changes in LDL-C would have suggested inhibition of 3-hydroxy-3-methylglutarylcoenzyme A reductase as a potential new therapeutic strategy By this reasoning, some of the more than 15 novel GWA loci discovered to date may harbor clinically useful drug targets and, thus, merit functional investigation.

Increasingly larger GWA studies with more than 100,000 participants of European descent (eg, by the Global Lipids Genetics Consortium), as well as GWA studies in other ethnic groups (eg, African Americans in the National Heart, Lung, and Blood Institute Candidate Gene Association Resource [NHLBI CARe]), are expected to uncover dozens more novel loci for which functional investigation will show numerous causal genes that will greatly enhance the understanding of lipoprotein metabolism and perhaps eventually lead to the development of new lipid-modifying medications.

Genome-wide Association Studies of Other Risk Factors for Myocardial Infarction

GWA studies have been performed for a number of cardio-vascular risk factors besides blood lipid concentrations. Studies on blood pressure have identified more than a dozen loci with common DNA variants that are associated significantly ($P < 5 \times 10^{-8}$) with systolic blood pressure or diastolic blood pressure (Table 4-2). ^{26,27} However, the effects of each SNP on blood pressure are quite small, in no case exceeding 1-mm Hg change per allele (see Table 4-2), and in most cases, potential functional links between the genes in each locus and the phenotype remain obscure.

One interesting exception is the chromosome 1p36 locus, which harbors five different genes with credible connections to blood pressure and cardiovascular disease: MHFTR, which encodes methylenetetrahydrofolate reductase, which catalyzes a critical step in homocysteine metabolism; CLCN6, which encodes a chloride channel; NPPA and NPPB, which encode atrial natriuretic peptide and B-type natriuretic peptide, respectively, which have vasodilatory effects; and AGTRAP, which encodes angiotensin II receptor-associated protein, a modulator of the renin-angiotensin-aldosterone axis. Although common DNA variants directly within the NPPA and NPPB genes have also been demonstrated to be highly associated with blood pressure, 28 it is difficult to know which of the five 1p36 genes (or combination of genes) exerts the effect on blood pressure detected by the GWA study; this lack of information highlights the general challenge that will be faced repeatedly by investigators seeking

TABLE 4—1	Loci Ass	ociated wit	h Blood Lipid Conce	entrations			
Unique Locus	LIVED	СН	SNP	Sample Size	P -Value	Gene(s) of Interest within or near Associated Interval	Associated Mendelian Lipid Disorders
1	LDL	1p13	rs12740374	19,648	2 x 10 -42	CELSR2-PSRC1-SORT1	-
2	LDL	2p24	rs515135	19,648	5 X 10 -29	APOB	Familial hypercholesterolemia
3	LDL	19q13	rs4420638	11,881	4 x 10 -27	APOE-APOC1-APOC4-APOC2	Type III hyperlipoproteinemia
4	LDL	19p13	rs6511720	19,648	2 x 10 -26	LDLR	Familial hypercholesterolemia
5	LDL	2p21	rs6544713	23,456	2 x 10 -20	ABCG5-ABCG8	Sitosterolemia
6	LDL	5q13	rs3846663	19,648	8 X 10 - 12	HMGCR	-
7	LDL	5q23	rs1501908	27,280	1 x 10 -11	TIMD4-HAVCR1	
8	LDL	20q12	rs6102059	28,895	4 x 10 -9	MAFB	
9	LDL	7p15	rs12670798	17,797	6 X 10 -9	DNAH11	
10	LDL	12q24	rs2650000	39,340	2 x 10 -s	HNF1A	-
11	LDL	1p32	rs11206510	19,629	4 x 10 -8	PCSK9	Familial hypercholesterolemia
12	HDL	16q13	rs1532624	21,412	9 x 10 -94	CETP	CETP deficiency
13	HDL	15q22	rs1532085	21,412	1 x 10 -35	Lipcsei	Hepatic lipase deficiency
14	HDL	16q22	rs2271293	21,412	8 x 10 - 16	CTCF-PRMT8-LCAT	LCAT deficiency
15	HDL	18q21	rs4939883	19,785	7 x 10 - 15	LIPG	-
16	HDL	9q31	rs3905000	21,412	9 x 10 - 13	ABCA1	Tangier disease
17	HDL	11p11	rs7395662	21,412	6 x 10 -11	MADD-FOLH1	-
18	HDL	9p22	rs471364	40,414	3 X 10 - 10	TTC39B	-
19	HDL	20q13	rs1800961	30,714	8 x 10 - 10	HNF4A	-
20	HDL	12q24	rs2338104	19,793	1 x 10 - 10	MMAB-MVK	-
21	HDL	19p13	rs2967605	35,151	1 x 10 -s	ANGPTL4	-
22	HDL	1q42	rs4846914	19,794	4 x 10 -s	GALNT2	-
23	TG	11q23	rs964184	19,840	4 x 10 -∞	APOA1-APOC3-APOA4-APOA5	Primary hypoalphalipoproteinemia
24	TG	8p21	rs12678919	19,840	2 x 10 -41	LPL	Familial hyperchylomicronemia
25	TG	2p23	rs1260326	19,840	2 X 10 -31	GCKR	-
26	TG	8q24	rs2954029	19,840	3 X 10 - 19	TRIB1	-
27	TG	7q11	rs714052	19,840	3 X 10 - 15	MLXIPL	-
28	TG	11q12	rs174547	38,846	2 X 10 -14	FADS1-FADS2-FADS3	-
29	TG	1p31	rs1167998	17,815	2 X 10 - 12	DOCK7-ANGPTL3	-
30	TG	19p13	rs17216525	19,840	4 x 10 -11	NCAN-CILP2-PBX4	-
31	TG	20q13	rs7679	38,561	7 x 10 -11	PLTP	-
32	TG	8p23	rs7819412	33,336	3 x 10 -8	XKR6-AMAC1L2	-

CETP, cholesteryl ester transfer protein; Chr, chromosome locus; HDL, high-density lipoprotein; LCAT, lecithin-cholesterol acyltransferase; LDL, low-density lipoprotein; SNP, single-nucleotide polymorphism; TG, triglycerides.

Data from Kathiresan S, Willer CJ, Peloso GM, et al: Common variants at 30 loci contribute to polygenic dyslipidemia, *Nat Genet* 41:56-65, 2009; and from Aulchenko YS, Ripatti S, Lindqvist I, et al: Loci influencing lipid levels and coronary heart disease risk in 16 European population cohorts, *Nat Genet* 41:47-55, 2009.

to understand the functional effects of GWA loci with multiple genes.

Type 2 diabetes mellitus is one of the most exhaustively studied phenotypes, having been analyzed in several successful phases of GWA studies of increasingly large size; to date, more than 20 genome-wide significantly associated loci have been identified. ²⁹⁻³⁴ Many of these loci harbor genes that appear to alter insulin processing and secretion by the pancreatic beta cell. For example, *TCF7L2* (transcription factor 7-like 2), the gene in the GWA locus most strongly associated with type 2 diabetes, encodes a transcription factor that interacts with the Wnt

signaling pathway and regulates proglucagon gene expression in gut endocrine cells 35 ; patients with diabetes risk-conferring variants in the TCF7L2 gene exhibit decreased levels of insulin secretion from beta cells. 36 Despite the fact that diabetes is a strong risk factor for cardiovascular disease, it remains unclear whether genes such as TCF7L2 that have been identified in diabetes GWA studies will prove to significantly contribute to cardiovascular disease.

TABLE 4-2	Loci Associated	d with Blood Pressur	e			
Unique Locus	Chr	SNP	<i>P</i> Value	Change in Blood Pressure (mm Hg) per Allele (SE)	Gene(s) of Interest within or near Associated Interval	Trait
1	10q24	rs11191548	7 x 10 ⁻²⁴	1.16 (0.12)	CYP17A1-NT5C2	SBP
2	15q24	rs1378942	1 X 10-23	0.43 (0.04)	CSK-ULK3	DBP
3	4q21	rs16998073	1 X 10-21	0.50 (0.05)	FGF5	DBP
4	12q24	rs653178	3 X 10-18	0.46 (0.05)	SH2B3-ATXN2	DBP
5	1p36	rs17367504	2 X 10-13	0.85 (0.11)	MTHFR-CLCN6-NPPA-NPPB-AGTRAP	SBP
6	12q21	rs2681492	4 x 10-11	0.85 (0.13)	ATP2B1	SBP
7	10q21	rs1530440	1 X 10 ⁻⁹	0.39 (0.06)	C10orf107	DBP
8	11p15	rs381815	2 x 10-₃	0.65 (0.11)	PLEKHA7	SBP
9	3p22	rs9815354	3 x 10∘	0.49 (0.08)	ULK4	DBP
10	17q21	rs16948048	5 x 10∘	0.31 (0.05)	ZNF652	DBP
11	17q21	rs12946454	1 x 10∘	0.57 (0.10)	PLCD3	SBP
12	10p12	rs11014166	1 x 10∘	0.37 (0.06)	CACNB2	DBP
13	12q24	rs2384550	4 x 10∘	0.35 (0.06)	TBX3-TBX5	DBP

Chr, chromosome locus; DBP, diastolic blood pressure; SE, standard error; SBP, systolic blood pressure; SNP, single-nucleotide polymorphism.

Data from Newton-Cheh C, Johnson T, Gateva V, et al: Genome-wide association study identifies eight loci associated with blood pressure, *Nat Genet* 41:666-676, 2009; and from Levy D, Ehret GB, Rice K, et al: Genome-wide association study of blood pressure and hypertension, *Nat Genet* 41:677-687, 2009.

Several non-traditional risk factors for cardiovascular disease have also been studied with GWA investigations. For example, C-reactive protein (CRP) and fibrinogen, two inflammatory biomarkers that are predictive of disease in prospective cohort studies, each have several loci that are significantly associated with the biomarker's blood concentration. ³⁷⁻⁴⁰ Not surprisingly, among the associated SNPs are variants in the *CRP* gene (for CRP) and in the *FGB* gene, which encodes fibrinogen beta chain (for fibrinogen). Also found to be associated with either of the two biomarkers were SNPs near a variety of metabolic, inflammatory, and immunity genes, which suggests that the blood biomarker levels integrate signals from multiple metabolic, inflammatory, and immune pathways.

Every clinical trait demonstrated to be associated with cardiovascular risk will probably be ultimately subjected to the GWA approach.

Genome-wide Association Studies of Myocardial Infarction and Coronary Artery Disease

Three GWA studies for coronary artery disease were published simultaneously in 2007: one from the Ottawa Heart Study, 41 one from the Icelandic company deCODE genetics, 42 and one from the Wellcome Trust Case-Control Consortium. 43 Despite using independent cohorts and different genotyping arrays, all three studies demonstrated the same novel locus on chromosome 9p21 to be associated with disease. Of particular note was the finding that genotypes of index SNPs in the 9p21 locus were not associated with any of the traditional risk factors for cardiovascular disease; this suggests that the genetic mechanism encoded in this locus operates through a previously unknown risk pathway. Furthermore, the mini mally defined locus (~ 58 kilobases in individuals of European descent) harbors no known genes, and so it is unclear how the causal DNA variant or variants in the locus influence phenotype. In subsequent studies, the association of the 9p21 locus with coronary artery disease and, specifically, MI has been replicated, as have a variety of other vascular phenotypes such as abdominal aortic aneurysm,

intracranial aneurysm, and peripheral arterial disease; these findings are suggestive of a pathogenetic mechanism in vascular tissue. $^{44-46}$

Besides the 9p21 locus, the study from the Wellcome Trust Case Control Consortium identified 43 SNPs in several additional loci associated with coronary artery disease at or near the statistical significance threshold of $P < 5 \times 10^{-8}$. A second set of GWA studies for either coronary artery disease or MI, each with several thousand disease cases, confirmed some of these loci and characterized several more associated loci. 45,47-49 To date, strong statistical evidence links more than a dozen loci to disease development (Table 4-3), and future GWA studies of larger size, such as those by the Coronary ARTery DISease Genome-wide Replication And Meta-analysis (CARDIoGRAM) consortium, are likely to identify more. Several of these loci are linked to blood lipid concentrations (see Table 4-3), but the remainder are not clearly associated with any of the traditional cardiovascular risk factors or even emerging biomarkers such as CRP. Functional characterization of these loci may reveal multiple risk pathways, previously unknown, that represent new therapeutic opportunities for the prevention of MI.

IMPLICATIONS OF GENETICS FOR CAUSALITY OF RISK FACTORS

The ability to perform genetic analyzes in large cohorts of individuals being monitored for incident cardiovascular events now makes it possible to test the relationships between cardiovascular risk factors and disease. Mendelian randomization is a technique in which DNA variants are used to address the question of whether an epidemiological association between a given risk factor and disease signifies a causal relationship between the two. ⁵⁰ If a DNA variant is known to directly influence an intermediate phenotype, and the intermediate phenotype is causal for disease, then the DNA variant should be associated with the disease to the extent predicted

TABLE 4-3	Loci Associa	ated with Myocardial Infarction or Coronar	y Artery Diseas	se		
Unique Locus	Chr	SNP	<i>P</i> Value	Odds Ratio (95% CI) per Risk Allele	Gene(s) of Interest within or near Associated Interval	Associated With Lipids?
1	9p21	rs4977574	3 x 10⋅4	1.29 (1.25-1.34)	CDKN2A-CDKN2B-ANRIL	No
2	1p13	rs646776	8 X 10-12	1.19 (1.13-1.26)	CELSR2-PSRC1-SORT1	Yes
3	21q22	rs9982601	6 X 10-11	1.20 (1.14-1.27)	SLC5A3-MRPS6-KCNE2	No
4	1q41	rs17465637	1 X 10-9	1.14 (1.10-1.19)	MIA3	No
5	10q11	rs1746048	7 X 10-9	1.17 (1.11-1.24)	CXCL12	No
6	6p24	rs12526453	1 X 10-9	1.12 (1.08-1.17)	PHACTR1	No
7	19p13	rs1122608	2 X 10-9	1.15 (1.10-1.20)	LDLR	Yes
8	2q33	rs6725887	1 x 10⋅8	1.17 (1.11-1.23)	WDR12	No
9	1p32	rs11206510	1 x 10∘	1.15 (1.10-1.21)	PCSK9	Yes
10	12q24	rs2259816	4 x 10⋅°	1.08 (1.05-1.11)	HNF1A	Yes
11	12q24	rs3184504	9 x 10∗	1.13 (1.08-1.18)	SH2B3	No
12	3q22	rs9818870	5 X 10 ⁻⁷	1.15 (1.11-1.19)	MRAS	No
13	6q26-6q27	rs2048327-rs3127599- rs7767084-rs10755578 (haplotype)	4 x 10-15	1.82 (1.57-2.12)	LPA	Yes

Chr, chromosome locus; CI, confidence interval; SNP, single-nucleotide polymorphism.

Data from Myocardial Infarction Genetics Consortium, Kathiresan S, Voight BF, et al: Genome-wide association of early-onset myocardial infarction with single nucleotide polymorphisms and copy number variants, *Nat Genet* 41:334-341, 2009; from Erdmann J, Grosshennig A, Braund PS, et al: New susceptibility locus for coronary artery disease on chromosome 3q22.3, *Nat Genet* 41:280-282, 2009; and from Tregouet DA, Konig IR, Erdmann J, et al: Genome-wide haplotype association study identifies the SLC22A3- LPAL2-LPA gene cluster as a risk locus for coronary artery disease, *Nat Genet* 41:283-285, 2009.

by (1) the effect of the DNA variant on the phenotype and (2) the effect of the phenotype on the risk of developing disease. Lack of the predicted association between the DNA variant and disease in an adequately powered sample would argue against a purely causal role for the intermediate phenotype in the pathogenesis of the disease.

This study design mimics a prospective randomized clinical trial, in which the randomization of each individual occurs at the moment of conception: genotypes of DNA variants are "assigned" to gametes in a random manner during meiosis, a process that is assumed not to be influenced by the typical confounders observed in observational epidemiologic studies; for example, a parent's disease status or socioeconomic status should not affect which of his or her two alleles of an SNP is passed to a child, each allele having an equal (50%) chance of being transmitted via the gamete to the zygote. In other words, Mendelian randomization should be unaffected by confounding or reverse causation. This technique has potential shortcomings: for example, it is only as reliable as the robustness of the estimates of the variant's effects on pheno type and effects of phenotype on disease, and the DNA variant is assumed not to influence the disease by means other than the intermediate phenotype being studied (pleiotropy), which in many cases may not be true. However, when this technique is carefully executed, the results can be as informative as those of a well-conducted randomized clinical trial.

Although no formal mendelian randomization studies of LDL-C and other lipid traits have yet been reported, studies in this vein have confirmed a causal relationship between LDL-C and cardiovascular disease. For example, nonsense coding variants in the *PCSK9* gene that were discovered in African Americans result in significantly reduced blood LDL-C concentrations; these reduced concentrations were, in turn, observed to be associated with the reduced incidence of CHD in a large African American cohort. ^{51,52} Similarly, a common missense coding variant in *PCSK9* in European Americans

associated with lower LDL-C levels was also found to be associated with a lower risk of CHD and MI. ^{52,53} More recently, 11 SNPs found to be associated with LDL-C in a GWA study were reported to be associated with CHD. ⁵⁴

In contrast, three independent mendelian randomization studies of variants in the *CRP* gene that affect blood CRP concentrations, performed in thousands of individuals, did not show an association between these variants and either ischemic vascular disease or CHD. ^{38,55,56} Although these findings cannot definitively rule out some causal role of CRP in MI, they suggest that any such causal role is minor in comparison with the role of LDL-C. They also suggest that the cardiovascular risk reduction obtained with rosuvastatin therapy in the Justification for the Use of Statins in Primary Prevention: an Intervention Trial Evaluating Rosuvastatin (JUPITER), ⁵⁷ in which patients with baseline normal LDL-C levels and elevated CRP levels were studied, resulted more from the lipid-lowering effects of the statin rather than its CRP-lowering effects.

A parallel line of evidence similarly casts doubt on the notion that inflammatory molecules such as CRP are critical mediators of cardiovascular disease. Of the 13 loci most highly associated with MI and coronary artery disease (see Table 4-3), 5 are related to blood lipid concentrations, which is strongly indicative of a causal relationship between lipid levels and disease. In contrast, none of the other 8 loci are clearly related to inflammation, which suggests that inflammatory molecules are of less pathobiological importance to MI than are lipid levels or, for that matter, to the as-of-yet-uncharacterized risk mechanisms represented by the 8 non lipid-related loci. This observation cannot be attributed to a bias of GWA studies against inflammatory gene SNPs, as much as classical inflammatory diseases such as rheumatoid arthritis and Crohn's disease have been found to be associated with numerous inflammatory gene SNPs at genome-wide significance. 58,59

Thus, although inflammation may contribute to the pathogenesis of MI, results of research with the currently available genetic techniques suggest that, on a population-wide basis, inflammation is of modest causal importance in comparison with other risk factors such as LDL-C.

UTILITY OF GENETIC RISK SCORES FOR DISEASE PREDICTION

Conventional cardiovascular risk algorithms such as the Framingham risk score, which includes several traditional risk factors and is generally limited to 10-year predictions, do not yield accurate predictions about many cardiovascular events. Much energy in the field of preventive cardiology has been directed towards the identification of novel risk factors that, when combined with conventional risk algorithms, will enable more accurate predictions of who will develop disease. In view of the partial heritability of cardiovascular disease, there is considerable interest in determining whether the use of genetic data will improve risk prediction.

A genetic risk score (ranging from 0 to 18) that accounts for nine SNPs associated with either LDL-C or HDL-C was found to be correlated with incidental cardiovascular disease in a prospective cohort study 60; each unfavorable allele (a single point in the score) conferred a 15% increase in risk after adjustment for traditional risk factors, including blood lipid concentrations. When stratified into groups with a high risk score or a low risk score, individuals with a high risk score were found to have an actual 63% increase in risk in comparison with those with a low risk score.

The association of the lipid genetic risk score with disease that was independent of blood lipid concentrations was attributed to the genetic risk score reflecting lifetime exposure to higher or lower lipid levels, whereas a single fasting lipid profile represents a snapshot of a patient's condition at the time the profile is measured. It is also possible that some of the lipid-associated SNPs have pleiotropic effects that contribute to cardiovascular disease but are not reflected in traditional risk factor measurements.

Addition of the genotype score to traditional risk factors did not significantly improve risk discrimination; no change was found in the C-statistic (area under the receiver operator characteristic curve). Nevertheless, modest numbers of individuals at intermediate cardiovascular risk, as judged by the Adult Treatment Panel III criteria, were correctly reclassified into a higher or lower risk category. Of note was that all of the lipid SNPs used in this genetic risk score predated the GWA studies reported since 2007; thus, the genetic risk score does not include dozens of SNPs now known to be associated with lipid levels. Those SNPs may be expected to significantly improve the predictive value of the risk score.

A comprehensive genetic risk score would include SNPs that are not associated with traditional risk factors—such as index SNPs in the chromosome 9p21 locus identified in GWA studies to be most highly associated with coronary artery disease and MI—and thereby have more independent predictive value than a lipid level-only genetic risk score. The 9p21 genotype by itself confers up to a 60% increase in risk in individuals with two unfavorable alleles. ^{41,45,61} A risk score that includes nine SNPs identified in GWA studies as being associated with early-onset MI, including an SNP at locus 9p21 and three SNPs associated with LDL-C, is even more highly associated with disease, with a 2.2-fold difference in risk for MI between extreme quintiles of risk score. ⁴⁵

Nevertheless, attempts to incorporate SNPs at locus 9p21 into risk-prediction models have yielded disappointing results to date. As with the lipid genetic risk score, adding the 9p21 genotype to traditional risk factors in prospective cohort studies

with men ⁶¹ and women ⁶² yielded no improvement in risk discrimination (as judged by C-statistic) and reclassified only small proportions of individuals to more accurate risk categories. However, investigators do await the evaluation of a comprehensive genotype score that includes many or all of the SNPs discovered to be strongly associated with cardiovascular disease.

Finally, as noted at the beginning of this chapter, a personal family history of early-onset MI in at least one parent more than doubles the risk of a cardiovascular event. As more genetic variants associated with disease are discovered, it will be important to assess whether a comprehensive genetic risk score will add any predictive value above and beyond simply asking about a patient's family history. For this reason, determining a genetic risk score may ultimately prove to be most useful in infants and children (whose parents may not be old enough to have developed coronary artery disease) for the purpose of determining lifetime cardiovascular risk and engaging in more stringent primary prevention practices.

UTILITY OF GENETICS FOR PERSONALIZED MEDICINE

Another potential use of genetics information is its application to pharmacogenetics: determining which individuals are more likely to benefit from (or to suffer an adverse effect from) the use of a particular medication. The design of pharmacogenetic studies is similar to that of traditional genetic studies except that the phenotype of interest, instead of being a disease or clinical trait, is the outcome upon receiving a therapy.

At least three examples of pharmacogenetic findings are relevant to the prevention or treatment of cardiovascular disease. First, the statin drugs are the most widely used medications used to lower lipid levels because of their consistent efficacy in reducing cardiovascular endpoints in numerous clinical trials. These trials have documented wide variability in individuals' response to statin therapy in the degree of LDL-C lowering. Pharmacogenetic studies in some of these trials, as well as in other cohorts, have reproducibly demonstrated that variants of SNPs in lipid level-related genes, HMGCR and APOE, are associated with the percentage decrease in blood LDL-C concentration experienced by statin users. 63-68 Thus, in principle, genotyping before initiation of lipid-lowering therapy could help predict response to statin drugs and guide practitioners in choosing among the statins (low- vs. high-potency) or choosing the starting dose for an individual patient - so-called personalized medicine.

A second, converse finding is that statin use occasionally causes myopathy that in extreme cases is life-threatening. A GWA study for statin-induced myopathy identified an SNP in the *SLCO1B1* gene as highly associated with this adverse effect. ⁶⁹ In individuals with two unfavorable alleles at this SNP, the risk of developing myopathy while they take statin therapy is 17 times higher than that in individuals with no unfavorable alleles. Thus, a genetic test for this SNP could be useful in screening patients before initiation of therapy, particularly if there is already concern that the patient is at risk for myopathy because of family history or has a personal history of muscle symptoms while receiving statin therapy. Patients with the risk-conferring genotype may wish to avoid statins and choose alternative therapies for lowering lipid levels.

The third example involves the antiplatelet agent clopidogrel, which is widely used in patients after acute coronary syndrome, percutaneous coronary intervention, or both. Clopidogrel is converted into its active metabolite by the CYP2C19 enzyme of hepatic cytochrome P-450. In three large

52 studies of patients receiving clopidogrel after acute coronary syndromes, individuals with reduced-function alleles of the *CYP2C19* gene experienced significantly higher rates of car diovascular death, myocardial infarction, and stroke. ⁷⁰⁻⁷² This is consistent with the finding in one of the studies that reduced-function allele carriers harbored lower plasma levels 4 of the active metabolite of clopidogrel. ⁷⁰ In principle, patients with reduced-function *CYP2C19* alleles would benefit from higher doses of clopidogrel or alternative antiplatelet medications such as prasugrel, although this remains to be tested in prospective clinical trials.

CONCLUSION

The development of the GWA technique has elucidated the genetics of cardiovascular disease and cardiovascular risk factors; studies with this technique have revealed numerous loci that represent previously unknown biological mechanisms and, ultimately, potential new therapeutic opportunities. Future studies from groups such as the Global Lipids Genetics -Consortium and CARDIoGRAM will extend these findings even further by screening very large populations and identify even more loci associated with blood lipid concentrations and coronary artery disease, and the NHLBÎ CARe and other studies will yield fresh insights into human genetics by applying GWA to non-European populations. Although it remains unclear whether genetics will be useful for cardio vascular risk prediction in adults, it may eventually be useful in other applications such as primary prevention and personalized medicine.

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CHAPTER 5

Novel Biomarkers and the Assessment of Cardiovascular Risk

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KEY POINTS

- Cardiovascular risk stratification must be improved, and biomarkers, genetic markers, and imaging provide the best avenue toward this improvement.
- All currently available markers (biomarkers and genetic markers) provide only limited to modest improvements in the ability to predict cardiovascular risk.
- In the future, the combination of genetic markers, imaging markers, and biomarkers will probably be used in an attempt to identify at-risk individuals while investigators continue to refine risk prediction with traditional risk factors.

The limitations of traditional coronary heart disease (CHD) risk stratification through the use of scores such as the Framingham Risk Score have been well documented and discussed. 1-3 The majority of individuals who have CHD events would have been classified as having low or intermediate risk by traditional risk stratification schemes, because most of the general population has low to intermediate 10-year (short-term) Furthermore, although the risk factors for CHD and stroke are similar, the risk prediction algorithms are different 4-9; therefore, an individual may have low risk for CHD and yet high risk for stroke, and vice versa. 10 In addition, although risk prediction tools are available, many clinicians do not use them, and those who do typically estimate only CHD risk and do not estimate risk for stroke, peripheral arterial disease, or heart failure. Newer tools that estimate total cardiovascular disease (CVD) risk are available 10 '1 and would be preferred to those that are limited to estimating CHD risk; however, the newer tools still focus on traditional risk factors and do not address longer term risk. Finally, most risk scores have been derived in populations with a predominance of one ethnicity, and the applicability of those scores to other ethnicities is therefore not known. Hence, improved CVD risk assessment tools are needed. Strategies to improve risk prediction have focused on identifying individuals who have an increased long term risk (ie, lifetime risk) 11 and in identifying novel markers. These additional markers include those identified on imaging, genetic markers, and biomarkers measured in plasma or urine.

CRITERIA FOR EVALUATING A NEW MARKER IN RISK PREDICTION

On average, more than 1100 reports of investigations of independent predictors or risk factors for various clinical outcomes are published every year, and CHD is one of the outcomes more frequently assessed. ¹² Some of the newly discovered markers have been reported to improve CHD risk prediction in comparison with traditional risk factors. Tzoulaki and colleagues assessed ¹³ studies

reporting improved CHD risk prediction beyond the Framingham risk score and found that the majority of the studies had design, analytical, or reporting flaws. A scientific statement from the American Heart Association ¹⁴ therefore recommends that certain important parameters be evaluated and reported to determine whether a marker adequately improves CHD risk prediction (Box 5-1).

Among the first things to consider is whether the marker is tested in an appropriate population. A cohort from a population based epidemiological study is ideal because the participants are representative of the population at large. Even in this cohort, however, there are limitations: for example, whether findings are generalizable to other ethnicities not studied. After basic analyses, including whether the marker is associated with the outcome of interest, odds ratio, risk ratio, and hazards ratio, the marker should be tested for (1) its ability to discriminate between persons who have the disease of interest (eg, CHD) and those who do not, (2) its accuracy in risk prediction, and (3) its effect on reclassifying individuals in the low- and intermediate-risk groups.

The ability of a marker to "discriminate" between persons with and those without a particular outcome is generally tested by describing the C-statistic, or the area under the receiver operating characteristic (ROC) curve, which essentially plots sensitivity against 1 specificity, or true-positive findings against true-negative findings. A value of 0.50 indicates that the marker has no more value than chance. However, the use of the Cstatistic in model selection (ie, to decide what variables to include in a model) has limitations. 15 Other tests based on likelihood, such as the likelihood ratio statistic or the Bayes information criterion, which adjusts for the number of variables in the model, are more sensitive 15,16 and may be better for use in model selection and as a measure of model fit. Another marker used in discrimination is the integrated discrimination improvement, which tests whether the novel marker correctly increases the predicted risk (ie, reclassification to a higher risk category) of persons who

2761

**BOX 5-1 Recommendations for Reporting **

- 1. Report the basic study design and outcomes in accordance with accepted standards for observational studies
- 2. Report levels of standard risk factors and the results of risk model, using these established factors
- Evaluate the novel marker in the population, and report:
 - a. Relative risk, odds ratio, or hazard ratio conveyed by the novel marker alone, with the associated confidence limits and P value
 - b. Relative risk, odds ratio, or hazard ratio for novel marker after statistical adjustment for established risk factors, with the associated confidence limits and P value
 - c. P value for addition of the novel marker to a model that contains the standard risk markers
- 4. Report the discrimination of the new marker:
 - a. C-index and its confidence limits for models with established risk markers
 - b. C-index and its confidence limits for models, including novel markers and established risk markers
 - c. Integrated discrimination index, discrimination slope, or binary R for the model with and without the novel risk marker
 - d. Graphic or tabular display of predicted risk in cases and noncases separately, before and after inclusion of the new marker
- 5. Report the accuracy of the new marker:
 - a. Display observed vs. expected event rates across the range of predicted risk for models without and with the novel risk marker
 - b. Using generally recognized risk thresholds, report the number of subjects reclassified and the event rates in the reclassified groups

From Hlatky MA, Greenland P, Arnett DK, et al: Criteria for evaluation of novel markers of

assess whether risk prediction is accurate. For this, a goodness-of-fit (LpPLA 2) level and amino-terminal pro B-type (or brain) natriuretic test is necessary to evaluate whether there is any difference between peptide (NT-proBNP) level. the predicted and observed risk. The number of individuals who are reclassified (ie, will change risk groups) by the inclusion of the risk marker of interest and the net effect of the reclassification (net reclassification index [NRI]) then needs to be determined. 17 The NRI, a statistical test designed to study the net effect of reactive protein was initially tested for association with CVD as reclassification, determines whether reclassifications were investigators increasingly appreciated the role played by appropriate; for example, if an individual was reclassified to a inflammation in the pathogenesis of atherosclerosis. 18 In several higher risk group and then had an event, the reclassification would studies, researchers have reported associations between hsCRP be considered appropriate ("good"), whereas if the individual was reclassified to a lower risk group and then had an event, the reclassification would be considered inappropriate ("bad"). The net evaluated the value of hsCRP in risk prediction in a number of effect of the "good" and "bad" reclassification determines the NRI, analyses. They first examined the value of hsCRP level when added and the clinical NRI is determined by the effect in the intermediaterisk group (in general, persons who have a 5% to 20% estimated 10- level, high-density lipopro tein cholesterol [HDL-C] level, smoking, year risk for CHD), in which the test might be used to refine risk and blood pressure) in the Women's Health Study. 29 In a cohort of assessment and need for treatment (Table 5-1).

It would be useful to show that a clinical strategy that used the events (116 myocardial infarctions, 217 coronary novel marker in risk prediction and in treating individuals can decrease the incidence of CHD. In this chapter, we discuss the use of biomarkers and genetic markers that have been studied for their use in the improvement of CVD risk prediction.

BIOMARKERS ASSESSED IN CARDIOVASCULAR **DISEASES RISK PREDICTION**

Several markers have been associated with CHD, stroke, or both, but only a very few have been tested for their influence on risk

Risk Category by		Risk (Category by Tradition Biomark		rs +
Iraditional Risk	Factors	<5%	5% to 20%	>20%	lotal
Individuals Who	Have a CI	inical Eve	nt <i>(n)</i>		
<5%		37	14	0	51
5% to 20%		5	85	16	106
>20%		0	4	24	28
Total		40	104	41	185
Individuals Who	Do Not Ha	ave a Clini	cal Event (n)		
<5%		1650	145	0	1795
5% to 20%		150	680	33	863
>20%		2	32	69	103

Calculation of Net Reclassification Index (NRI)

cardiovascular risk; a scientific statement from the American Heart Association. Circulation 119:2408-2416, 2009.

857

102

1802

NRI = [(number of individuals with events among those who were reclassified to a higher risk group/total number of individuals with events) — (number of individuals with events among those who were reclassified to a lower risk group/total number of individuals with events)] - [(number of individuals without events among those who were reclassified to a higher risk group/total number of individuals without events) — (number of individuals without events among those who were reclassified to a lower risk group/total number of individuals without events)].

For the data in this table:

Total

TABLE 5-1

- 1. Number of individuals with events among those who were reclassified to a higher risk group/total number of individuals with events = (14 + 16)/185 = 30/185 = 0.162
- 2. Number of individuals with events among those who were reclassified to a lower risk group/total number of individuals with events = (5 + 4)/185 = 9/185 = 0.049
- 3. Number of individuals without events among those who were reclassified to a higher risk group/total number of individuals without events = (145 + 33)/2761 = 188/2761 =
- 4. Number of individuals without events among those who were reclassified to a lower risk group/total number of individuals without events = (150 + 2 + 32)/2761 = 174/2761 = 0.066

Therefore, NRI = (0.162 - 0.049) - (0.064 - 0.066) = 0.113 - 0.005 = 0.115, or 11.5%

have the event and decreases the predicted risk of those who do not. prediction. The marker that has been best studied is high-sensitivity C-reactive protein (hsCRP) level. Other markers that appear Although these tests of discrimination are important, they do not promising include lipoprotein-associated phospholipase A 2

C-Reactive Protein

C-reactive protein is a nonspecific marker of inflammation. Clevel and incidental CHD, stroke, or both. 19-28

In view of the consistent association, Ridker and associates 29 to variables used in the Framingham risk score (age, total cholesterol 15,048 women aged 45 and older, 390 women had incident CVD

56 revascularization procedures, 65 deaths from cardiovascular causes, Reynolds risk score was associated with an NRI of 5.3% and a and 100 ischemic strokes) in an average follow-up period of 10 years. clinical NRI of 14.2%. Other analyzes have also suggested that the Although adding hsCRP level to a risk prediction model based on NRI for adding hsCRP level is approximately 5% to 7%. 33,34 Framingham variables only marginally improved the area under the However, in a case-control study of individuals in the European ROC curve (to 0.815, in comparison with 0.813 for the model without Prospective Investigation into Cancer and Nutrition (EPIC)-Norfolk hsCRP level), 5 other tests of discrimination, such as the Bayes study, the NRI for adding hsCRP level was 12.0%. 35 information criterion, suggested that a model that included hsCRP level would be better. According to model calibration tested with the Hosmer- hsCRP levels was studied in the Justification for the Use of Statins Lemeshow goodness-of-fit test, the model with hsCRP level was a better in Primary Prevention: an Intervention Trial Evaluating fit when expected and observed events were compared. Of the Rosuvastatin (JUPITER). Individuals with low-density lipoprotein individuals predicted to have a 5% to 20% risk over 10 years, about 20% cholesterol (LDL-C) levels lower than 130 mg/dL and hsCRP levels were reclassified after the addition of hsCRP level.

prediction could be improved with the inclusion of several novel adverse cardiovascular events, and the trial was discontinued early markers (eg, levels of hsCRP, hemoglobin A1c, homo cysteine, because of clear benefit. 36 Yang and coworkers 37 analyzed data on soluble intercellular adhesion molecule-1, apolipo proteins) that had participants in the Atherosclerosis Risk in Communities (ARIC) been identified since the Framingham risk score had been described. study according to the entry criteria for JUPITER; their findings They divided the Women's Health Study cohort into a model suggested that elevated hsCRP level confers high risk regardless of derivation cohort (n = 16,400) and a model validation cohort (n = LDL-C levels (either < 130 mg/dL or > 130 mg/dL) and after various 8158). The variables that resulted in the best fitting model included traditional risk factors are taken into account. age, hemoglobin A1c in subjects with diabetes, current smoking, lipoprotein(a) levels (if apolipoprotein B level > 100 mg/dL), Preventive Services Task Force (USPSTF) 38 concluded that there is apolipoprotein B level, apolipoprotein AI level, parental history of strong evidence that hsCRP level is associated with incident CHD, myocardial infarction (at age < 60 years), and natural logarithms of moderate evidence that hsCRP level can help in risk stratification of systolic blood pressure and hsCRP level. Ridker and colleagues then the intermediate-risk group, but insufficient evidence that reducing simplified this model for clinical use by substituting levels of total hsCRP level can prevent CHD events. However, in its systematic cholesterol and HDL-C for levels of apolipoproteins B-100 and AI review of nine "emerging" CHD risk factors, including hsCRP level, and eliminating the measurement of lipoprotein(a) level (Table 5-2). the USPSTF concluded that current evidence does not support the This Reynolds risk score, 31 which differed from the Framingham use of any of these factors in further risk stratification. 39 Similarly, risk score mainly in its use of hsCRP level and parental history of other investigators have questioned whether adding hsCRP level myocardial infarction, was found to have better model has any additional value in risk stratification. 40 Part of the reason discrimination and calibration and reclassified 40% to 50% of that these questions have been raised is the significant correlation of individuals in the intermediate-risk group into higher risk or lower hsCRP level with traditional risk factors and its minimal effect on risk categories. However, no patient was reclassified from the low- the area under the ROC curve. risk group (<5% CHD risk over 10 years) to the high-risk group (> 20% CHD risk over 10 years) or vice versa; this suggests that prior Survey (NHANES) data, Miller and associates reported that hsCRP probability of disease should be considered in determining for levels were rarely high (> 3 mg/L) in the absence of traditional risk whom additional testing is recommended.

Best-Fitting Model	Clinically Simplified Model: Reynolds Risk Score
Age	Age
Systolic blood pressure	Systolic blood pressure
Current smoking	Current smoking
hsCRP	hsCRP
Parental history of MI < age 60	Parental history of MI < age 60
Hemoglobin A1c (if diabetic)	Hemoglobin A1c (if diabetic)
Аро В-100	Total cholesterol
About Al	HDL-C

Notes: The Reynolds Risk Score was originally described in women and has since been described in men by means of the same clinically simplified model. 33

Apo AI, apolipoprotein AI; apo B-100, apolipoprotein B-100; HDL-C, high-density lipoprotein cholesterol; hsCRP, high-sensitivity C-reactive protein; Lp(a), lipoprotein(a); MI, myocardial

From Ridker PM, Buring JE, Rifai N, et al: Development and validation of improved algorithms for the assessment of global cardiovascular risk in women: the Reynolds Risk Score, JAMA 297:611-619, 2007.

The Reynolds risk score was subsequently described in men as well: in comparison with a traditional model, the Reynolds risk score reclassified 18% of subjects in the Physicians Health Study II, including 20% of subjects at intermediate risk, and was associated with a better model fit and discrimination. 32 In addition, the

More recently, a strategy of treating individuals with elevated of 2 mg/L or higher were treated with rosuvastatin; treatment with Ridker and colleagues 30 then investigated whether risk this drug was associated with a 44% relative risk reduction in major

The 2009 evaluation of hsCRP level by the United States

In an analysis of National Health and Nutrition Examination factors associated with CHD, occurring in 4.4% of men and 10.3% of women, and that elevations in hsCRP levels that were attributable to a borderline or abnormal CHD risk factor occurred in 78% of men and 67% of women. 41 Epidemiological studies such as the Framingham Heart Study and the ARIC study have also demonstrated that the effect of hsCRP level on improving the area under the ROC curve is minimal and not statistically significant. 42,43 However, using area under the ROC curve as the only metric to evaluate value in risk stratification can be suboptimal, because the C-statistic is based solely on ranks and is not as sensitive as measures based on likelihood. In fact, several well-established risk factors such as LDL-C and HDL-C may add little to the area under the ROC curve when added to other traditional risk factors. 15

Our own impression of the available data is that hsCRP level can help identify higher risk individuals among those classified as having intermediate short-term (10-year) risk for CHD by traditional risk prediction algorithms. However, it is unclear whether hsCRP level is a risk marker or a risk factor; that is, it is unclear whether hsCRP level plays a role in the pathogenesis of atherosclerosis or adverse cardiovascular events, or whether it is merely a bystander marking other changes that lead to atherogenesis and adverse cardiovascular events

events. Genetic studies have identified several loci associated with risk prediction may be better. Additional studies are needed to hsCRP levels but not with CVD, 44,45 which suggests that hsCRP examine whether pharmacological treatment of patients who have levels may be a risk marker. However, whether it is a risk marker or elevated LpPLA 2 levels can reduce CVD events. LpPLA 2 level may a risk factor should not affect the ability of hsCRP level to predict be a risk factor, not only a risk marker, and a large outcomes trial is risk.

associated with increased risk for CHD and stroke. In our opinion, and identify the role for LpPLA 2 level in CVD risk stratification. a clinically relevant number of individuals are reclassified, and a prospective trial has shown that treatment of individuals who have Amino-Terminal Pro-B-Type Natriuretic Peptide elevated hsCRP levels, "normal" LDL-C levels, and intermediate CHD risk can reduce both CHD and stroke. In addition, an expert panel convened by the National Academy of Clinical Biochemistry concluded, on the basis of a thorough literature review for a number for acceptance for risk assessment in primary prevention. 46

Lipoprotein-Associated Phospholipase A 2

LpPLA 2 level is another biomarker that has consistently been shown to be associated with both CHD and stroke. ^{22,47-51} LpPLA 2, which is predominantly associated with LDL in the circulation, is thought to mediate its inflammatory effects through its action on oxidized expression of adhesion molecules. 52

LpPLA 2 level has been evaluated as a marker for improving risk prediction. In the ARIC study, LpPLA 2 level was the only marker asymptomatic elderly participants were associated with increased asymptomatic elderly participants were associated with increased asymptomatic elderly participants. (of 19 markers studied, including hsCRP level) that significantly risk for CVD death and total mortality rate, and participants with increased the area under the ROC curve (by 0.006) when added to elevations of both markers had even higher risk. traditional risk factors that included age, race, sex, total cholesterol level, HDL-C level, systolic blood pressure, antihypertensive Other Markers medication use, smoking status, and diabetes. 43 However, in a more recent report from the EPIC-Norfolk study in which several markers were examined for their ability to improve risk prediction when added to a Framingham risk score-based model, only hsCRP level improved the C-statistic significantly; LpPLA 2 level had no significant effect. 35 Addition of LpPLA 2 level in this study resulted in an NRI of 1.7% and a clinical NRI of 8.8%, whereas adding hsCRP level was associated with an NRI of 12.0% and a clinical NRI of 28.4%. However, the model fit was better with LpPLA 2 level than with hsCRP level.

In view of the strong association of LpPLA 2 level with stroke (ischemic), Nambi and colleagues, using an analysis of a case-cohort random sample (n = 949, of whom 183 had incident ischemic stroke) from the ARIC study, evaluated whether LpPLA 2 level could improve stroke risk prediction. 10 Nambi and colleagues classified individuals' 5-year risk for stroke as low (< 2%), intermediate (2% to 5%), or high (> 5%) on the basis of a traditional risk factor model that included age, sex, race, current smoking, systolic blood pressure, LDL-C level, HDL-C level, diabetes, antihypertensive medication, and body mass index and then added hsCRP and LpPLA 2 levels separately and together to the analysis. Overall, adding LpPLA 2 level significantly improved the area under the ROC curve (from 0.732 to 0.752; 95% confidence interval [CI] for change in area under the ROC curve, 0.0028 to 0.0310), whereas adding hsCRP level did not significantly increase the area under the ROC curve (from 0.732 to 0.743; 95% CI for change in area under the ROC curve, - 0.0005 to 0.0183). However, adding both LpPLA 2 and hsCRP levels, as well as their interaction, resulted in the best improvement in the area under the ROC curve, which increased to 0.774 (95% CI for change in area under the ROC curve, 0.0182 to 0.0607). The addition of hsCRP level, LpPLA 2 level, and their interaction reclassified 4%, 39%, and 34% of the individuals originally classified as being at low, intermediate, and high risk, respectively.

In summary, LpPLA 2 level has not been as well studied as hsCRP level, especially with regard to improving risk prediction. Available data suggest that its ability to improve CHD risk prediction may be modest, but its ability to improve ischemic stroke

examining whether inhibition of LpPLA 2 in patients at high risk can In summary, there is consensus that elevation in hsCRP level is reduce CVD events. 52a Further studies will be needed to evaluate

B-type (or brain) natriuretic peptide (BNP) is a cardiac hormone secreted by cardiomyocytes in response to pressure and ventricular volume overload. The amino-terminal fragment of its prohormone (NT-proBNP), which has traditionally been thought of as a marker (NT-proBNP), which has traditionally been thought of as a marker of emerging risk factors, that only hsCRP level met all the criteria for congestive heart failure, has also been associated with both CHD and stroke. 53 The contribution of NT-proBNP level in risk stratification was examined in the Rotterdam study, 54 in which NTproBNP level was analyzed with traditional risk factors to investigate its ability to predict 10-year risk of CVD. For a group of 5063 individuals older than 55 years and free of CHD, addition of NT-proBNP level to traditional risk factors significantly improved the C-statistic both in men (0.661 to 0.694; change in C-statistic, 2 phospholipids, releasing lyso-phosphatidylcholine and oxidized 0.033; 95% CI, 0.012 to 0.052) and in women (0.729 to 0.761; change Ω nonesterified fatty acids, both of which are capable of attracting in C-statistic, 0.032; 95% CI, 0.016 to 0.047) and resulted in an NRI monocytes to an atherosclerotic lesion and further inducing the of 9.2% (95% CI, 3.5% to 14.9%; P = 0.001) in men and 13.3% (95% CI of 9.2% (95% CI, 5.5% to 14.5%, I = 0.001) in women. In the Rancho Bernardo Study, $\frac{8}{8}$ 54a increased NT-proBNP levels or detectable troponin T levels in

Several other markers also have associations with CVD; however, information regarding their use in CVD risk stratification is limited. In the analysis from the ARIC study noted previously, in which researchers examined the effect of adding various markers (n = 19) to traditional risk factors, only LpPLA 2 level improved the area under the ROC curve. 43 Rana and associates 35 investigated the effect of adding levels of hsCRP, myeloperoxidase, LpPLA 2, secretory phospholipase A 2 group IIA (sPLA 2), fibrinogen, paraoxonase, macrophage chemoat tractant protein-1 (MCP-1), and adiponectin to analyzes of CHD risk stratification. Overall, hsCRP level was the only marker that significantly improved the area under the ROC curve (to 0.65, from 0.59 for a Framingham risk score-based model; P = 0.005). Level of hsCRP was also associated with the best NRI and clinical NRI (12% and 28.4%, respectively), and sPLA $_{\rm 2}$ level was the next best (6.4% and 16.3%, respectively). However, when model fit was examined, adding hsCRP or paraoxonase or MCP-1 level to the Framingham risk score was associated with lack of model fit, whereas the addition of the other markers was associated with a good model fit. In the intermediate-risk group, the greatest numbers of individuals were accurately reclassified with the addition of sPLA 2 level, followed by levels of fibrinogen, LpPLA 2, adiponectin, and myeloperoxidase. In separate case-control analyzes from the

Multiple Markers

but not their use in risk stratification (reviewed by Koenig 55).

was 0.70 in a model that included age, sex, and the multimarker provided additive prognostic value over LDL-C level. 59a score; 0.76 in a model with age, sex, and conventional risk factors; and 0.77 in a model with all predictors.

Melander and associates 57 evaluated the additional value of 6 biomarkers (levels of hsCRP, cystatin C, LpPLA ₂ , midregional proadrenomedullin [MR-proADM], midregional pro- **FOR CORONARY HEART DISEASE** atrial natriuretic peptide, and NT-proBNP) in 5067 participants without CVD from Malmo, Sweden (mean age, 58 years). After Numerous new discoveries have helped investigators link genetic statistic for prediction of CHD events (increase in C-statistic, 0.007; P = 0.04), whereas NT-BNP and MR-proADM levels best improvement was not statistically significant (increase in Cstatistic, 0.009; P = 0.08). Very few individuals were reclassified: (Table 5-3) are described in this section. 8% of the study population was reclassified for CVD risk prediction and 5% for CHD risk prediction. Similarly, Genetic Variation in the Human Genome improvements in NRI for CVD and CHD were nonsignificant, The human genome comprises millions of DNA base pairs that although improvements in clinical NRI were significant (7% and 15%, respectively, largely through reclassification to a lower risk category).

Multiple markers have also been studied in older individuals. In one study in individuals older than 85 years, traditional risk factors were poor predictors of cardiovascular mortality, and of the markers studied (levels of hsCRP, homo cysteine, folic acid, and interleukin-6), homocysteine level was the best predictor of cardiovascular mortality (area under the ROC curve, 0.65; 95% CI, 0.55 to 0.75). On the other hand, with the use of biomarkers (levels of troponin I, NT-proBNP, cystatin C, and hsCRP); the C-statistic improved from 0.664 for traditional risk factors alone to 0.766 (difference, 0.102; 95% CI, 0.056 to 0.147) in the whole cohort and from 0.688 to 0.748 (difference, 0.059; 95% CI, 0.007 to 0.112) in subjects without (26%, P = 0.005). Overall, this study was limited by the fact that only 136 subjects died from CVD.

Hence, even with the use of multiple markers, a consistently reliable set of markers has not been identified for CVD risk prediction. Of the novel markers studied, the addition of BNP level to hsCRP level appears the most reliable.

Assessment of apolipoprotein B concentration and measurement of lipoprotein particle sizes with nuclear magnetic resonance (NMR) have been suggested as tests that may refine and improve risk Because many of these markers improve risk prediction prediction in comparison with cholesterol measurements currently marginally, efforts have been made to evaluate the value of a used clinically. Mora and colleagues 59 examined the association of multimarker approach by combining several biomarkers. With these tests with CVD and their ability to improve risk prediction in many of these multimarker approaches, the researchers examined the Women's Health Study, a study of healthy female health care primarily the association of markers (in concert) with CHD/CVD professionals aged 45 years or older. Although both NMR lipid profile and apolipo protein B concentration were associated with Wang and coworkers 56 assessed 10 biomarkers (levels of CVD after adjustment for nonlipid risk factors, the hazard ratios hsCRP, BNP, N-terminal pro-atrial natriuretic peptide, were similar to those for traditional lipid measures. The C-index was aldosterone, renin, fibrinogen, D-dimer, plasminogen- 0.784 for the model with nonlipid risk factors and ratio of total activator inhibitor type 1, and homocysteine, and the urinary cholesterol to HDL-C levels, and it was not significantly different albumin-to-creatinine ratio) in the Framingham Heart Study (with the addition of LDL level measured by NMR (0.785) or n = 3209) for their ability to predict major adverse apolipoprotein B level (0.786). NRI also did not show net cardiovascular events. BNP level (hazard ratio = 1.25) and improvement; in comparison with nonlipid risk factors and the total urinary albumin-to-creatinine ratio (hazard ratio = 1.20) had cholesterol-to-HDL-C ratio, NRI was 0% with NMR-measured LDL the strongest association with major adverse cardiovascular level and 1.9% with apolipoprotein B. This finding suggests that events, and BNP level (hazard ratio = 1.40), hsCRP level these novel lipid measures do not significantly enhance risk (hazard ratio = 1.39), and urinary albumin-to-creatinine ratio prediction in comparison with the traditional lipid measure of total (hazard ratio = 1.22) had the strongest association with death, cholesterol-to-HDL-C ratio. However, other studies in populations but none of the markers affected the C-statistically with higher baseline triglyceride values have demonstrated that significantly. The C-statistic for major cardiovascular events apolipoprotein B level and other measures of LDL particle number

GENETIC MARKERS AND ASSESSMENT OF RISK

using a backwards elimination model to identify the best variants to human disease processes. Genetic and epidemiological markers for prediction of CVD events (n = 418) and CHD studies of cardiovascular genetics and CHD in particular have events (n = 230) (median follow-up, 12.8 years), they reported identified genetic variants directly associated with CHD and CHD that hsCRP and NT-proBNP levels best improved the C- risk factors. However, the practical clinical implementation of this information for management and prevention of CHD continues to be evaluated. The major studies in which researchers have improved prediction of CVD events, although the evaluated the application of genetic variants associated with CHD in risk prediction and preventive cardiovascular management

constitute either coding regions, which code for proteins that are essential for cell function, or noncoding regions of unknown significance. One of the major characteristics of the human genome is its interindividual variation. This variation in genomic content and structure between individuals is large, and its importance in normal function varies. There are rare variants with a large effect on disease risk, common variants that usually have a small effect on disease susceptibility, and variants with no apparent influence on known disease.

The most frequent type of genomic sequence variation in the Zethelius and associates 58 reported significant improvement in human genome is the single nucleotide polymorphism (SNP). A prediction of CHD death in individuals older than 75 years SNP is a change in a single base pair at a specific genomic locus, so that the same single base pair is not in that locus for everyone; there may be a different base pair in a subgroup of the population. It is estimated that there is 1 SNP every 1000 base pairs and about 3 million base pair differences between any given two human genomes. Some of these SNPs are inherited together as part of a CVD. The NRI for adding all the biomarkers was significant block of DNA called a *haplotype*. This phenomenon is useful in research because it enables a single SNP (a "tag" SNP) to be tested as a marker

TABLE 5-3	Statistic	Statistical Metrics for Examining the Clinical Utility of Genetic Variants to Improve CHD Risk Prediction								
Subjects, n	Gender	Variant(s)	I Traditional Risk Factor Model	Improvement in Area unde the ROC Curve	er NRI	CNRI	Reference			
2742	Men	9p21	FRS	0.02	«4.3%*		Talmud et al ⁷²			
22,000	Women	9p21	FRS	0	2.70%	«7%*	Paynter et al ⁷⁰			
9998	Both	9p21	ACRS	0.004	0.80%	6.20%	Brautbar et al ⁷¹			
5414	Both	GRS	FRS	0	«6.1%*		Kathiresan et al ⁷⁴			

^{*}Estimated on the basis of the published reclassification table.

ACRS, ARIC [Atherosclerosis Risk in Communities] Risk Score; CHD, coronary heart disease; CNRI, clinical net reclassification index; FRS, Framingham risk score; NRI, net reclassification index.

genome and their interindividual variation, they are natural adjustment for traditional risk factors (including age, smoking, candidates for research on differences in disease susceptibility systolic blood pressure, cholesterol level, and HDL-C level) was 1.70 between individuals. According to the "common disease-common (95% CI, 1.19 to 2.41) for individuals who are homozygous for the variant" hypothesis, 59b complex diseases such as atherosclerosis and risk allele. Adjustment for family history modestly decreased the CHD are caused by not one gene but rather multiple genes, each of hazard ratio, which was suggestive of some correlation between the which contributes a small additive effect towards a certain threshold 9p21 risk allele and family history. However, there was no that results in the overall condition. SNPs are the ideal tool with statistically significant association between the two (P = 0.48). which to examine and discover genes or noncoding areas that contribute to diseases such as CHD.

different approaches have been applied, including the candidate- to 0.64, the increase was not statistically significant. Calibration, gene approach, which had limited success, and genome-wide examined with the Hosmer-Lemeshow metrics, revealed a association (GWA) studies, which have successfully identified nonsignificant P value, indicating a good fit for the models with and multiple loci associated with various disease conditions and traits. without the 9p21 risk allele. After the addition of the 9p21 risk allele, GWA studies are based on the testing of thousands and up to a approximately 22% of individuals were reclassified. NRI and million SNPs at once to identify loci associated with a disease, such clinical NRI were not calculated, but 63% of patients reclassified as CHD, and traits, such as LDL-C level. Important considerations were assigned to a more accurate risk as reflected by a more for both candidate-gene and GWA approaches are the need to appropriate event rate in their new category. Additional metrics correct the statistical metrics used for discovery for multiple testing, assessing model fit – the likelihood ratio and the Bayes information replication of the results in a study population that is similar to the criterion—were improved by the addition of the 9p21 risk allele. original discovery cohort, and examining the association in other populations and ethnicities.

The 9p21 Chromosomal Region and Coronary Heart and frequency as the 9p21 risk allele) to a traditional risk factor-Disease

In 2007, two independent GWA studies reported a number of SNPs in a 58-kilobase interval on the 9p21 chromosomal region that demonstrated a strong association with CHD in white persons. 60,61 These SNPs defined a single haplotype (ie, they were closely linked together and inherited together) and were found to be associated with increases in CHD risk of approximately 20% in heterozygotes and approximately 40% in homozygotes. After the initial report, multiple studies replicated and validated this association in the white population 62-64 and demonstrated the association in additional populations, including Han Chinese, 65 East Asian, 66 South Korean, ⁶⁷ Hispanic, ⁶⁶ and Italian. ⁶⁸ However, this association tested for association with fraditional risk factors, the 9p21 risk allele was not demonstrated in African American populations. 60,66 Interestingly , there are no known genes in this 58-kilobase interval addition of the 9p21 risk allele did not significantly in the 9p21 chromosomal region, although two genes, CDKN2A and CDKN2B, are located adjacent to it. Results of one study suggested that the 9p21 risk allele has a major role in the cardiac expression of CDKN2A and CDKN2B, which directly affect the proliferation properties of vascular cells. 69

The importance of the 9p21 chromosomal region was not only its association with CHD but was also the high frequency of the risk allele in the white population: 45% to 55% in various studies. 62,63,70-⁷² The combination of a large effect size with high population frequency made the 9p21 risk allele an attractive marker with which to enhance CHD risk prediction.

Talmud and associates 72 were the first investigators to evaluate whether the addition of the 9p21 risk allele to traditional risk factors

0.004 0.80% 6.20% Brautbar et alm 0 «6.1%* Kathiresan et alm Biomarkers and the Biomarker for multiple SNPs. The less frequent SNP or allele usually has a improves CHD risk prediction. To examine their hypothesis, they frequency of greater than 5%, which is defined as the minor allele used the Northwick Park Heart Study II (NPHS-II), a prospective Because of the relatively large numbers of SNPs in the human population sustained 270 CHD events. The hazard ratio after

Discrimination was examined by adding the 9p21 risk allele to a model with age and clinical practice (site of patient recruitment) To identify SNPs that may be associated with disease processes, only. Although the area under the ROC curve increased from 0.62 \overline{\text{\text{\text{\text{P}}}}}

> Because of the lack of improvement in discrimination, Talmud and associates 72 suggested a potential approach to enhance the use of the 9p21 risk allele to improve risk prediction. Incremental addition of 1 to 10 hypothetical variants (with similar effect sizes based model significantly improved the area under the ROC curve after the addition of the first variant. These findings suggested that SNPs may be combined to construct a genetic risk score that may improve risk prediction; this concept is developed later in this section.

> Following the analysis by Talmud and associates, 72 who used a cohort comprising only men, Paynter and colleagues 70 examined the clinical utility of the 9p21 risk allele added to traditional risk factors in a cohort of 22,129 white women who were prospectively monitored for a period of approximately 10 years; in this cohort, 715 total incident CVD events (CHD and stroke) were sustained. The hazard ratio was 1.15 per 9p21 risk allele for CVD events, and when had a modest association with family history and diabetes. The

60 increase the C-index in comparison with models based on the Framingham risk score and Reynolds risk score. Only 2.7% of the women were reclassified, most of whom (approximately 86%) were reclassified correctly. The NRI and integrated discrimination improvement were 2.7% and 0.001, respectively; clinical NRI was not calculated. Goodness of fit 5 was tested with the Hosmer-Lemeshow metrics, which demonstrated a good fit for both models, with and without the 9p21 risk allele. The conclusion of Paynter and colleagues was that the addition of the 9p21 risk allele to traditional risk factors in analysis was not useful clinically. However, because this large cohort had a relatively low number of incident events (715 CVD events for 22,129 subjects), the statistical power of the study was substantially limited.

Brautbar and coworkers 71 examined whether the addition of the 9p21 risk allele to traditional risk factors improves CHD utility of adding the 9p21 risk allele to traditional risk factors SNPs capture that value. along with the Framingham risk score. The ACRS is based on respectively). Goodness of fit examined with the Gronnesby- through a known intermediate phenotype. Borgan metrics was better for the model with the 9p21 risk allele, although both models did not demonstrate a good fit.

clinical utility of reclassification, Brautbar and coworkers clinical utility for CHD risk prediction. measured the baseline LDL-C distribution in the categories for 5% to 10% and 10% to 20% CHD risk over 10 years before reclassification. In both risk groups, approximately 90% had Assessment of Risk for Cardiovascular Disease LDL-C levels higher than 100 mg/dL, and thus reclassification to a higher risk category would have practical implications for many individuals by changing LDL-C target goal and initiation level for lipid-modifying therapy based on National reclassification.

Genetic Risk Score

extensive efforts were made to identify a panel of SNPs that prevention of CHD. would enable better estimation of CHD risk. The first study to examine this question included approximately 15,000 individuals from the ARIC study who developed approximately 1400 CHD events. 73 SNPs to be examined were chosen on the basis of prior GWA and candidate-gene studies. After extensive effort to genotype the SNPs in ARIC, frequency and association with CHD were tested in both African American and white subjects. Within each race, SNPs

associated with P values higher than 0.10 were excluded, which left 11 SNPs for each race. The SNPs were then modeled for an additive effect of the risk-raising allele and were individually evaluated by Cox proportional hazards models. The genetic risk score, comprising these 11 SNPs, was added to traditional risk factors on the basis of the ACRS model. Calculation of the area under the ROC curve demonstrated modest improvement for African American subjects (0.758 to 0.769) and marginal improvement for white subjects (0.764 to 0.766). Reclassification was not examined in this study. The study's conclusion was that the improvement in discrimination was modest and the evidence was not clinically applicable. However, the additive approach taken in this study was widely adopted in subsequent genetic studies of CHD risk prediction.

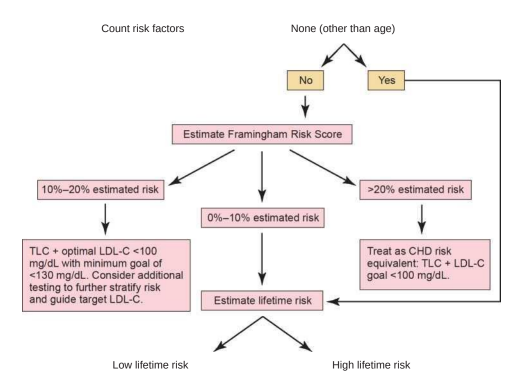
An interesting approach to constructing a genetic risk score for risk prediction in the ARIC study. The ARIC population - CHD prediction was presented by Kathiresan and associates, 74 who examined included 9998 white middle-aged men and women examined the additive effect of SNPs that were already known to be who were monitored for approximately 14 years, of whom associated with LDL-C and HDL-C. The main hypothesis was that 13.5% had 1349 incident CHD events. The calculated hazard although these SNPs were associated with an intermediate ratio was 1.2 per allele for the 9p21 risk allele after adjustment phenotype that is already well established for CHD risk assessment, for traditional risk factors. The ARIC Cardiovascular Risk Score they represent a measurement of lifelong exposure to that particular (ACRS), a model based on traditional risk factors that was intermediate phenotype. This exposure has an additional predictive created and tested in the ARIC study, was used to evaluate the value beyond that of the intermediate phenotype itself, and these

To examine this hypothesis, the investigators used a cohort of age, gender, smoking, diabetes, systolic blood pressure, 5414 subjects who developed 238 CHD events. When discrimination antihypertensive medication use, total cholesterol level, and was examined, the C-statistic was the same for the models with and HDL-C level. The frequency of the risk allele was 49% in the without the genetic risk score. After reclassification, the NRI was entire cohort and, as expected, significantly higher among modestly improved. However, reclassification was poor, especially subjects with CHD events. Discrimination was evaluated by for the 10% to 20% CHD risk category. In our opinion, both calculation of the area under the ROC curve, which was discrimination and reclassification in this study did not show a modestly but significantly improved for the model with the substantial improvement, which suggests that it would be more 9p21 risk allele over the model without it (0.780 and 0.776, beneficial to use SNPs that are associated directly with CHD and not

Paynter and colleagues 75 examined the hypothesis suggested by Kathiresan and associates 74 in the Women's Genome Health Study. When the 9p21 risk allele was added to traditional risk SNPs were included in a genetic risk score based on a literature factors, approximately 13% in the intermediate-low category search for GWA studies. The genetic risk score was based on 101 (5% to 10% risk over 10 years) and intermediate-high category SNPs known to be associated with an intermediate phenotype of (10% to 20% risk over 10 years) were reclassified in both the CHD. A model based on Adult Treatment Panel III variables with ACRS and Framingham risk score models. The NRI and clinical and without the genetic risk score showed no improvement in the NRI after the addition of the 9p21 risk allele were 0.8% and C-statistic and no significant increase in the NRI (0.5). These results 6.2%, respectively, for the ACRS model. The clinical NRI for the further suggest that genetic risk score models based on SNPs that Framingham risk score model was 6.8%. To evaluate the are not associated with an intermediate phenotype may have better

Summary on the Use of Genetic Markers in

The use of genetic information for CHD risk prediction is an attractive possibility and has made considerable progress. However, no clinical guidelines currently exist for either the conduct of GWA Cholesterol Education Program Adult Treatment Panel III studies or the assessment of SNPs to refine the assessment of CVD guidelines. In summary, the largest effect on reclassification in risk. Multiple examinations of genetic risk scores as means to this analysis was on the intermediate -risk categories, 90% of improve risk prediction have demonstrated only limited clinical whom had LDL-C levels above the recommended goals after utility. However, the discovery of the 9p21 risk allele has demonstrated that certain genetic markers with large effect size and high population frequency may help improve risk prediction models in the future. As the area of discovery of genetic markers develops, better genetic risk prediction models are possible and may Before the discovery of the 9p21 chromosomal region, lead to the use of genetic markers to improve risk prediction and



TLC + optimal LDL-C <130 mg/dL. Consider treatment for LDL-C >160 mg/dL.

Further risk stratification with hsCRP, CIMT, coronary calcium score, or genetic risk score:

- If reclassified to higher risk groups, LDL-C goal <130 mg/dL, with an optional goal <100 mg/dL.
- If no reclassification, optimal LDL-C <130 mg/dL; treat LDL-C >160 mg/dL.

Continue periodic, recommended risk factor screening

FIGURE 5-1 Risk assessment algorithm. CHD, coronary heart disease; CIMT, carotid intima-media thickness (measured by ultrasound); hsCRP, high-

TLC + optimal LDL-C <100 mg/dL with minimum goal of <130 mg/dL $\,$

sensitivity C-reactive protein; LDL-C, low-density lipoprotein cholesterol; TLC, therapeutic lifestyle change. (From Nambi V Ballantyne CM: "Risky business': ten years is not a lifetime, Circulation 119:362-364, 2009.)

CONCLUSION

Cardiovascular risk stratification must be improved, and biomarkers, genetic markers, and imaging provide the best avenue towards this improvement (Figure 5-1). ⁷⁶ However, all currently available markers (biomarkers and genetic markers) confer only limited to modest improvements in the ability to predict risk. The combination of genetic markers, imaging markers, and biomarkers will probably be used in concert in an attempt to identify at-risk individuals while investigators continue to refine risk prediction with traditional risk factors. However, both the clinical utility and cost-effectiveness of such approaches need to be determined. On the other hand, risk stratification will be of clinical use (for disease management) primarily when treatment plans change with assessed risk or when therapies are available to target novel risk factors. Currently, the only risk factors whose goals vary according to estimated risk are LDL-C level and blood pressure. With the price of statins decreasing, the cost- and risk benefit ratios may allow a single cut point, as for other risk factors, in determining the need for statin therapy in the future. Hence, while risk stratification needs continued improvement, simultaneous advances in therapeutics are also crucial for reaching the eventual goals of personalized cardiovascular risk stratification and primary prevention.

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CHAPTER 6

Advanced Risk Assessment in Patients with Kidney and Inflammatory Diseases

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KEY POINTS

- The number of patients around the world with chronic kidney disease (CKD) has increased alarmingly.
- Even mild to moderate worsening of kidney function has become an independent risk factor for cardiovascular morbidity and mortality.
- In the future, novel risk markers—such as cystatin C, adiponectin, and possibly new inflammatory markers other than C-reactive protein (CRP) and microalbuminuria—may be used to assess risk for cardiovascular events.
- Management of patients with CKD is important for protection against both progression of kidney disease and progression of cardiovascular disease.
- Close monitoring and follow-up are key in the therapeutic management of patients with CKD.

Chronic kidney disease (CKD), defined as persistent kidney damage reflected by a glomerular filtration rate (GFR) of less than 60 mL/min/1.73 m ² for 3 months, ¹ is a major public health problem worldwide. More than 8 million people in the United States have stage 3 CKD, and the number is rising. This trend in CKD is reflected around the world, not just in the United States. ²

Patients with stage 3 or higher CKD have higher rates of cardiovascular morbidity, manifested by higher incidences of heart myocardial arrhythmias. failure. and infarctions . Progression of CKD in these patients to end-stage kidney disease - defined as a GFR of less than 10 mL/min/1.73 m² further increases the risk for cardiovascular events; the annual mortality rate has improved since 2000 but remains approximately 19% per year. ³ In earlier stage nephropathy (ie, GFR > $60 \text{ and } < 90 \text{ mL/min}/1.73 \text{ m}^{\frac{1}{2}}$), less is known regarding cardiovascular risk. 4-6 Investigators increasingly appreciate, however, the fact that risk markers such as microalbuminuria that are associated with vascular inflammation are indicative of higher cardiovascular risk (Table 6-1).7

PATHOPHYSIOLOGY

abnormalities are commonly observed in patients with CKD that may enhance their risk for cardiovascular disease (CVD) events. Although the precise mechanism by which CKD increases CVD risk is not fully elucidated, most cases of CKD are clearly associated with increased oxidative stress and magnified inflammatory responses at the level of the vasculature. Endothelial dysfunction is an early event in people with CKD, and microalbuminuria is associated with the presence of endothelial dysfunction. ⁷ In most patients with advanced CKD, atherosclerosis is accelerated characterized by more advanced, heavily calcified plaques that extend to both the intima and medial layers of the coronary vessels. 8,9 Increased expression of several cytokines, as well as of macrophages, plays a role in the evolution of the plaque development.

Several inflammatory markers have been implicated potential triggers atherosclerotic complications. Highsensitivity C-reactive protein (hs-CRP) is considered a biomarker of chronic systemic inflammation, as well as a mediator of atherosclerosis . Of patients with end-stage kidney disease, 20% to 50% have been shown have elevated CRP levels. Hyperhomocysteinemia is also a predictor of future CVD events in patients with established coronary artery disease and in patients with type 2 diabetes, as well as those undergoing dialysis. 12,13 Adiponectin level also plays an important role in modulating atherosclerosis and is decreased in people with impaired glucose homeostasis or diabetes. In the Mild and Moderate Kidney Disease (MMKD) study group, patients with low adiponectin levels experienced significant cardiovascular events.

The amount of nitric oxide present is reflective of how well the endothelium is functioning. It has a protective role in that it inhibits vascular muscle cell proliferation, platelet aggregability, and the adhesion of monocytes to the endothelium. The enzyme responsible for the genesis of nitric oxide can be inhibited by endogenous methyl arginine production such as asymmetric dimethylarginine (ADMA). Levels of ADMA are postulated to be increased in patients with advanced nephropathy, and such elevation is recognized as a putative biomarker in cardiovascular and kidney disease. 15 Two large clinical trials, the Coronary Artery Risk Determination investigated in the Influence of ADMA Concentration (CARDIAC) and the AtheroGene Study, demonstrated that ADMA factor independent risk cardiovascular disease. In these studies, baseline ADMA levels were independently predictive of cardio vascular events. 16,17

Hypertension is a complex phenotype because neurohumoral factors such as angiotensin II, norepinephrine, and other cytokines, as well as chronic volume overload, exert inflammatory and growth promoting effects in the cardiovascular system. ¹⁸ Angiotensin II is a proinflammatory substance and a recognized growth

TABLE 6-1

Risk Factors and Novel Risk Markers of Chronic Kidney
Disease in Predicting Cardiovascular Morbidity and Mortality

Risk Factors for Cardiovascular Disease

Older age Race Smoking history Hypertension Diabetes

Diabetes
Asymmetric dimethylarginine dyslipidemia
Anemia
Chronic kidney disease

Novel Risk Markers for Cardiovascular Disease

Cystatin C level C-reactive protein level Microalbuminuria Adiponectin level Albumin level

promoter. The sympathetic system not only is a major regulator of cardiovascular function but also affects immune response, as does angiotensin II. It is interesting to note that in patients with CKD, circulating levels of norepinephrine are directly related to the muscular component of the left ventricle. Norepinephrine levels are also a strong and independent predictor of death from cardiovascular causes. Chronic volume overload is a major stressor, and in the long run, the deleterious effects of volume overload depend on the fact that hemodynamic burden activates a series of adaptive processes that modify the very structure of the myocardium. ¹⁸

The aforementioned factors—together with retention of toxins, increased calcium intake, and decreased phosphate excretion; abnormalities in bone mineral metabolism; and poor nutrition state—all increase inflammatory markers and potentiate vascular disease. 14,19-21

RISK IN COMMUNITY-BASED POPULATIONS

CKD itself is a major risk factor for cardiovascular events. In many cohort studies, risk for coronary heart disease (CHD) has been assessed in relation to changes in CKD stage. The findings of these studies have led to the formation of recommendations from both The National Kidney Foundation and the American College of Cardiology/American Heart Association that CKD be considered as a CHD risk equivalent. Many physicians may not be aware that increases in risk for CHD parallel reductions in GFR, highest risk being at GFR values lower than 45 mL/min. ⁴

The Framingham Heart Study ²² is one prospective, community -based study of the burden of CVD in patients with kidney disease. The study has revealed that the majority of patients with mild to moderate CKD are older, are more likely to be obese, have lower levels of high-density lipoprotein (HDL), and higher triglyceride levels. They also have a high prevalence of hypertension, diabetes, and elevated levels of low-density lipoprotein (LDL). The CKD population is less likely to achieve optimal control of blood pressure and controlled hemoglobin A1c concentrations of less than 7%.

Another large cohort study involved the Kaiser Permanente Renal Registry. Among 1,120,295 adults within a large, integrated system of health care delivery, the GFR was estimated. ⁴ After adjustment, the risk of death increased as the estimated GFR decreased below 60 mL/min/1.73 m². The adjusted hazard ratio for cardiovascular events also increased inversely with the estimated GFR. The adjusted risk of hospitalization with a reduced estimated GFR followed a similar pattern. The findings highlighted the clinical and public health importance of CKD. In the Atherosclerosis Risk in Communities (ARIC) study, ²³ participants with a GFR of 15 to 59 mL/min/1.73 m ² (hazard ratio, 1.38; 95% confidence interval, 1.02 to 1.87) and 60 to 89 mL/min/1.73 m ² (hazard ratio, 1.16; 95% confidence interval, 1.00 to 1.34) had an increased adjusted risk for atherosclerotic CVD events, in comparison with subjects with normal GFR

levels, after a mean follow-up of 6.2 years.

In the National Health and Nutrition Examination Survey (NHANES), rates of CVD-related mortality were 4.1, 8.6, and 20.5 deaths per 1000 person-years among participants with estimated GFRs of higher than 90, 70 to 89, and lower than 70 mL/min/1.73 m², respectively. Those with an estimated GFR lower than 70 mL/min/1.73 m² had significantly higher relatively risks of 6 death from cardiovascular disease (1.7; 95% confidence interval, 1.3 to 2.1) and from all causes. ²⁴

CARDIOVASCULAR RISK FACTORS IN PATIENTS WITH CHRONIC KIDNEY DISEASE

CVD is the major cause of morbidity and mortality among patients with CKD. Most patients share risk factors, including diabetes, hypertension, obesity, lipid abnormalities, and smoking (see Table 6-1). Even people with early stage 3 nephropathy (ie, estimated GFR < 60 mL/min/1.73 m 2) have a higher risk of mortality than those with a GFR above 60 mL/min/1.73 m 2 .

Diabetes is the most common cause of CKD, accounting for nearly 50% of all new cases of renal replacement therapy. In a cohort of Chinese patients with type 2 diabetes who did not have macrovascular disease or end-stage renal disease, all-cause mortality increased from 1.2% to 18.3% as kidney function deteriorated from stage 1 to stage 4. ²⁵ Hypertension is another modifiable risk for CVD. The degree and duration of hypertension strongly influence outcomes and also accelerate CKD progression. ²⁶ Most patients with CKD have both hypertension and diabetes as comorbid conditions, and the effect on CVD risk is more than additive.

Obesity is a major global health concern and may precede the development of many CVD risk factors, including diabetes, hypertension, and dyslipidemia. In the Framingham Heart Study, ²⁷ obesity was noted to be associated with increased risk of developing stage 3 CKD during nearly 20 years of follow-up. This finding suggests that the association of obesity with stage 3 CKD may be mediated by vascular disease risk factors.

Patients with CKD are at high risk for insulin resistance and other features of the classical metabolic syndrome. The association of higher body mass index (BMI), insulin resistance, hyperglycemia, and hypertriglyceridemia supports the notion that early in the disease state, other well-known CVD risk factors are present and may be magnified by the presence of advanced CKD. 9

Overweight and obesity are also associated with increased risk of proteinuria and risk of worsening kidney function, inasmuch as increased levels of proteinuria are associated with faster CKD progression. ²⁸ Conversely, in patients undergoing dialysis, the relationship is different: The greater the BMI with better nutrition, the lower the incidence of CVD events. In short, survival is higher in patients with end-stage kidney disease who have higher BMIs.

The sequelae of CKD, such as anemia and low active vitamin D levels, may also contribute to the increased risk for CVD. 29,30 In a registry cohort study of 5549 adults hospitalized with acute myocardial infarction or unstable angina, pro found anemia was independently associated with increased mortality rate (hazard ratio, 1.8 for hemoglobin levels of < 9 vs. > 12 g/dL) among patients with an estimated GFR of 30 to 59. 31

6

Γrial	Design	Population	Sample Size	Intervention	Duration	Outcome
Pravastatin Pooling Pr (PPP)	Post hoc subanalysis of West of Scotland Coronary Prevention Study (WOSCOPS), Cholesterol and Recurrent Events (CARE), and Long-term Intervention with Pravastatin in Ischaemic Disease (LIPID)	Moderate CKD (CG-GFR, 30-59 mL/ min/1.73 min:	4491	Pravastatin, 40 mg/day	«5 years	Decile risk (HR, 0.77) of adjusted incidence of primary outcome
Heart Protection Study (HPS)	Post hoc subanalysis		1329	Simvastatin, 40 mg/day	5 years	Decile risk (HR, 0.72) of major vascular events
		CKD (serum creatinine clearance >110 ^mol/L for women and >130 ^mol/L for men but <200 gmol/L)				
inglo-Scandinavian Cardiac Outcomes Trial-Lipid Lowerin Arm (ASCOT-LLA)	g	Renal dysfunction (microalbuminuria, proteinuria)	6571	Atorvastatin, 10 mg/day	3.3 years	Decile risk (HR, 0.61) of primary endpoint
eterans' Affairs High Density Lipoprotei Intervention Trial (VA-HIT)	n	Predialysis CKD	1046	Gemfibrozil, 1200 mg/ day) 5.3	Decile risk (HR, 0.73) of primary outcome and decile risk (HR, 0.74) of combined outcome of coronary death, nonfata MI, or stroke
Study of Heart and Re Protection (SHARI		Predialysis status Hemodialysis Peritoneal dialysis	9000 (6000 before dialysis and 3000 undergoing dialysis)	Simvastatin, 20 mg, and ezetimibe, 10 mg	4 years	Major vascular events, rates progression to ESR in patients before dialysis

CG-GFR, glomerular filtration rate calculated with Cockcroft-Gault equation; CKD, chronic kidney disease; ESR, end-stage renal disease; HR, hazard ratio; MI, myocardial infarction; RCT, randomized controlled trial.

Elevated cholesterol levels are very prevalent among people with estimated GFR values lower than 60 mL/min/ 1.73 m ². Lowering cholesterol levels in people who have diabetes and an estimated GFR higher than 60 mL/min/1.73 m ² is beneficial, as observed in the Scandinavian Simvastatin Survival Study (4S), in which patients with type 2 diabetes had a 2.5-fold greater risk for coronary artery disease than did nondiabetic patients. ^{32,33} Ongoing trials are currently being conducted to examine the benefit of lowering cholesterol levels in early- to moderate-stage CKD ³⁴; however, it is clear that lowering cholesterol levels in patients with advanced-stage CKD who are undergoing dialysis does not alter CVD outcomes. ^{35,36}Thus, early use of statins slows nephropathy progression and reduces CVD risk, whereas late use once dialysis has been instituted fails to alter CVD risk (Table 6-2).

There is also a relationship between cardiovascular disease and microalbuminuria. ³⁷ In a secondary analysis of the Multiple Risk Factor Interventional Trial, the presence of minimal proteinuria conferred nearly a 2.5-fold greater risk for cardiovascular morbidity events. ³⁸ The presence of microalbuminuria in patients with diabetes is a useful marker for patients at greatest risk for the development of macrovascular disease. ⁷ In addition, post hoc analyzes from the Losartan Intervention For Endpoint reduction in hypertension (LIFE) trial clearly demonstrated that reduction in albuminuria progression over time is associated with a lower incidence of CVD outcomes.

ASSOCIATION OF CHRONIC KIDNEY DISEASE AFTER MYOCARDIAL INFARCTION

There is a significant rise in mortality among CKD patients after an acute coronary event. The prognosis of patients after acute myocardial infarction may be poor partly because of a relatively of presentations, resulting from increased number underdiagnosis and undertreatment. 40,41 For example, the presence of dyspnea in a patient with end-stage renal disease may be mistakenly attributed to volume overload. In a survey, 44% of patients undergoing dialysis present with chest pains, in comparison with 68% of patients with CKD who are not undergoing dialysis. 40 Medications are underused, and aggressive therapy such as thrombolysis and angiography is not prescribed because of further increase of creatinine levels, which leads to acute renal failure. As a result, the rate of 1-year mortality after an acute myocardial infarction in patients with CKD is approximately 50%. 42 The Coopera tive Cardiovascular Project performed a cohort study with 130,099 elderly patients with mild to moderate CKD who had myocardial infarctions. 43 The rates of 1-year mortality were 46% among patients with stage 3 CKD (creatinine level, 1.5 to 2.4 mg/dL) and 66% among patients with stage 4 CKD (creatinine level, 2.5 to 3.9 mg/dL). Problems with the management of these types of patients arise because

they received less therapy-whether aspirin, beta blockers, thrombolytic therapy, angiography, or angioplasty-during

In another retrospective cohort study, outcomes after an acute myocardial infarction were compared between patients with The number of CKD patients around the world has increased varying levels of CKD and patients without CKD. In hospital mortality rates were 2% among patients with normal kidney advanced CKD, 21% among those with severe kidney failure, and adiponectin, and possibly new inflammatory markets cannot 30% among those with end-stage kidney disease. Postdischarge hs-CRP and microalbumin uria may be used to assess risk for some control of patients who received acute reperfusion cardiovascular events. Management of patients with CKD is a control of patients with CKD is a control of patients with CKD is a control of patients. function, 6% among those with CKD, 14% among those with and mortality. In the future, novel risk markers such as cystatin C, therapy (odds ratio, 0.7), aspirin (odds ratio, 0.7), and beta blocker important for protection against both progression of kidney disease therapy (odds ratio, 0.7). 44

MEASURES TO PREVENT CARDIOVASCULAR **EVENTS IN PATIENTS WITH CHRONIC KIDNEY DISEASE**

Risk modification and lifestyle changes can decrease cardiac events in patients with CKD. In general, similar conditions apply to patients who have CKD and those who do not with regard to smoking cessation, maintaining ideal body weight, active lifestyle, and glycemic control in diabetes.

The treatment of hypertension is important in patients with CKD to protect against both progressive kidney disease and cardiovascular disease. As stated in the guidelines of the Joint National Committee on Prevention, Detection, Evaluation, and Treatment of High Blood Pressure, ⁴⁵ the goal for blood pressure in proteinuric CKD is lower than 130/80 mm Hg to slow the rate of 9 progression of kidney disease. Angiotensin converting enzyme inhibitors and angiotensin receptor blockers have been shown to 10. Grootendorst DC, de Jager DJ, Brandenburg VM, et al: Excellent agreement between C-reactive significantly reduce cardiovascular morbidity and mortality in multiple, large, prospective randomized trials. 46 They have a dosedependent beneficial effect on atherosclerosis progression and may prevent the development and recurrence of atrial fibrillation. The 12. Heinz J, Kropf S, Luley C, et al: Homocysteine as a risk factor for cardiovascular disease in Perindopril Protection against Recurrent Stroke Study (PROG RESS) had a post hoc analysis in which 29% of patients with creatinine clearance of less than 60 mL/min were evaluated; the use of perindopril, an antihypertensive therapy, reduced the risk of all cardiovascular events in patients with CKD. 47

The use of lipid-lowering agents such as statins, as previously discussed, is very useful in slowing nephropathy and reducing CVD risk among patients with an estimated GFR above 30 mL/min, but these agents were not useful in altering CVD events in patients undergoing dialysis (see Table 6-2). The 2003 Kidney Disease 17. Schnabel R, Blankenberg S, Lubos E, et al: Asymmetric dimethylarginine and the risk of Outcomes Quality Initiative (K/DOQI) guidelines recommend a goal LDL cholesterol level of less than 100 mg/dL. 48 Hypertriglyceridemia and low HDL concentrations are common lipoprotein abnormalities in patients with CKD. Fibrates effectively 19. Horl WH, Cohen JJ, Harrington JT, et al: Atherosclerosis and uremic solute retention. Kidney Int lower triglycerides and elevate HDL-cholesterol concentrations, which could complement the effectiveness of statins. The Veterans' 20 . Affairs High Density Lipoprotein Interventional Trial (VA-HIT) demonstrated lower cardiovascular events with gemfibrozil in patients with creatinine clearance of less than 75 mL/min. 49

is sparse. Three studies have shown the benefit of giving aspirin to patients with CKD. A retrospective observational analysis from the Dialysis Outcomes and Practice Patterns Study (DOPPS) revealed 24. Muntner P, He J, Hamm L, et al: Renal insufficiency and subsequent death resulting from that aspirin resulted in a decreased risk of stroke (relative risk, 0.82) in all patients undergoing dialysis. ⁵⁰ A secondary subgroup analysis of the Hypertension Optimal Treatment (HOT) study found that in patients with serum creatinine levels higher than 1.3 mg/dL, lowdose aspirin (75 mg/day) significantly reduced the numbers of cardiovascular events and myocardial infarctions. 51 The safety of aspirin was evaluated in the first 67 United Kingdom Heart and Renal Protection (UK-HARP I) study. 52 It was not associated with an increased risk of major bleeding in comparison with placebo. The National Kidney Foundation suggests that the prescription of lowdose aspirin is probably safe in most patients with CKD. ¹

CONCLUSION

alarmingly. Even mild to moderate worsening of kidney function has become an independent risk factor for cardio vascular morbidity and progression of cardiovascular disease. Close monitoring and follow-up are key in the therapeutic management of patients with CKD.

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SECTION ||

Atherothrombosis and Antiplatelet Therapy

CHAPTER 7

KEANTIPLATE THE TANK

- Six randomized trials of aspiring therapy for the primary prevention of cardiovascular disease have demonstrated a 12% relative reduction in the risk of major adverse
- In large randomized trials of secondary prevention, aspirin has resulted in a 25% reduction in serious vascular events.

cardiovascular events.

- For a decision of whether to initiate aspirin in a primary prevention setting, current U.S. Preventive Services Task Force guidelines recommend incorporating an estimation of an individual patient's risk of hemorrhage.
- Dual-antiplatelet therapy with clopidogrel and aspirin is the mainstay of treatment after acute coronary syndromes and percutaneous coronary intervention.
- Clopidogrel resistance is an increasingly recognized phenomenon that underscores the importance of newer antiplatelet agents such as prasugrel and the oral P2Y₁₂ receptor antagonist ticagrelor.
- Novel agents targeting the platelet P2Y₁₂ and thrombin receptors are currently being studied in phase II and III trials and hold promise for the future.

Platelet activation plays a central role in the development of atherothrombosis, and antiplatelet therapy is thus a cornerstone of prevention and treatment of cardiovascular disease. Initial platelet activation and rapid platelet amplification occurs in response to potent agonists such as thromboxane A 2, adenosine diphosphate (ADP), and thrombin. Investigators' understanding of these pathways has led to the development of pivotal pharmacotherapies for treating cardiovascular disease. For example, the thromboxane inhibitor aspirin has resulted in substantial reductions in cardiovascular morbidity, and some authors have estimated that it could prevent 100,000 vascular deaths per year. ² In this chapter, we review the mechanism of action, data from primary and secondary prevention trials, and guidelines for antiplatelet agents currently in widespread use. We also discuss ongoing trials of novel antiplatelet agents directed at platelet targets such as the ADP receptor and the less exploited thrombin receptor.

ASPIRIN

Mechanism of Action

Acetylsalicylic acid, or aspirin, is the most widely used antiplatelet agent in the treatment of cardiovascular disease. Aspirin exerts its main antiplatelet effect by acetylating a serine residue on the cyclo oxygenase (COX) or prostaglandin H synthase enzyme and thus irreversibly inhibiting the action of this

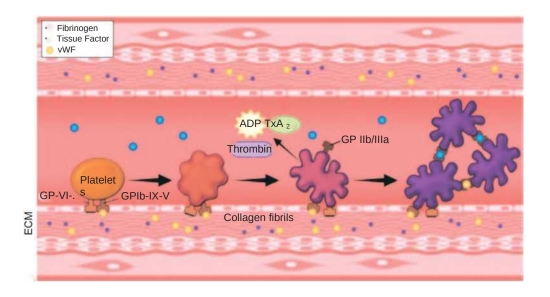
enzyme. ³ After exposure to aspirin, the anucleated platelet is largely unable to synthesize COX during its 7- to 10-day lifespan. ⁴COX enzymes, which exist in at least two iso forms, are responsible for the production of prostaglandins and thromboxane from ara chidonic acid. Preferential inhibition of COX-1 results in decreased production of thromboxane A ₂ , a potent mediator of platelet aggregation. ⁵ Other potential mechanisms of action include inhibition of intrinsic nitric oxide synthase ⁶ and inhibition of transcription factors involved in inflammation ⁷ (Figure 7-1).

Secondary Prevention

The salutary effect of aspirin for the second year prevention of cardiovascular disease is well established. In the first small studies to examine this relationship in patients with a history of myocardial infarction, the results were suggestive of a mortality benefit but were statistically inconclusive. 8-10 More convincing evidence arose from the Anti platelet Trialists' Collaboration (ATC), a metaanalysis of 31 randomized trials of antiplatelet therapy primarily with aspirin in patients who had sustained prior myocardial infarction, stroke, transient ischemic attack (TIA), or unstable angina. 11 Of 29,000 patients, those platelet treated with anti therapy demonstrated a 25% reduction in the odds of suffering a recurrent vascular event. 11 In a second study, the ATC demonstrated an 18% reduction in the odds of vascular death among patients at high risk, as defined by history of myocardial infarction, stroke, TIA, or unstable angina. 12

Although intuited from smaller randomized studies, ¹³ the benefit of aspirin in the setting of an acute myocardial infarction was persuasively demonstrated in the Second International Study of Infarct Sur vival (ISIS-2). ¹⁴ In this trial of 17,187 patients with a suspected acute myocardial infarction, a 162.5-mg daily dose of aspirin administered for 1 month significantly





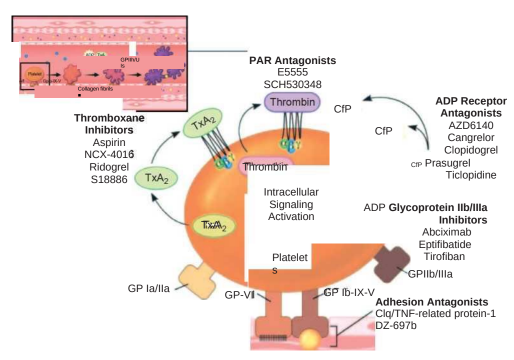


FIGURE 7-1 Platelet activation and the mechanism of thrombus formation. **A**, Endothelial injury exposes components of the extracellular environment such as collagen and von Willebrand factor (vWF). After binding to these components by means of glycoprotein receptors, platelets adhere to the subendothelium and become activated. Activation of the platelet causes a conformational change in the shape of the platelet, release of adenosine diphosphate (ADP) and thromboxane A 2 (TxA 2), and formation of thrombin on the platelet surface. The release of factors such as ADP and TxA 2 causes activation of circulating platelets and amplifies the platelet response. These responses cause the platelet glycoprotein (GP) llb/llla receptor to change shape and increase its affinity for adhesive proteins such as vWF and fibrinogen. Platelet aggregation ensues, and the additional interaction of the platelet aggregate with thrombin and fibrin results in thrombus formation. **B**, The agonists ADP, TxA 2, and thrombin bind to G protein-coupled receptors and trigger an intracellular signaling cascade. Several antiplatelet therapies are directed at inhibiting the interaction between these agonists and their respective receptors such as the ADP receptor antagonists, thromboxane inhibitors, and emerging protease activating receptor (PAR) antagonists. C1q/TNF, C1q complex/tumor necrosis factor; P2Y 1 and P2Y 12, G protein-coupled purinergic receptors. (From Meadows TA, Bhatt DL: Clinical aspects of platelet inhibitors and thrombus formation, Circ Research 100:1261-1275, 2007.)

reduced early vascular mortality in comparison with placebo (9.4% versus 11.8%, respectively). $^{14}\,\rm The$ protection afforded by aspirin extended to patients with unstable angina in a study of 1266 male veterans. $^{15}\,\rm In$ this randomized, placebo-controlled trial, a daily 324-mg buffered aspirin administered for 12 weeks resulted in a 51% reduction in myocardial infarction or death. $^{15}\,\rm Similar$ results emerged from the study by the Research Group on Instability in Coronary Artery Disease (RISC), which demonstrated a 57% to 69% reduction in the rate of the combined endpoint of myocardial infarction

TABLE 7—1 Primary Prevention Trials of Aspirin

Trial	Year of Publication	Population	Aspirin Dose	Stroke	Relative Risk Myocardial Infarction	All-Cause Mortality
British Doctors' Trial	1988	5139 male physicians	500 mg or 300 mg daily	1.13	0.97	0.89
Physicians' Health Study	1989	22,071 male physicians	325 mg every other day	1.22	0.59	0.96
Thrombosis Prevention Trial	1998	5085 men	75 mg daily	0.98	0.68	1.06
Hypertension Optimal Treatment Study	1998	18,790 men and women	75 mg daily	0.98	0.64	0.93
Primary Prevention Project	2001	4495 men and women	100 mg daily	0.67	0.69	0.81
Women's Health Study	2005	39,876 female health professionals	100 mg every other day	0.83	1.02	0.95

Adapted from Meadows T, Bhatt DL: Clinical aspects of platelet inhibitors and thrombus formation, Circ Res 100:1261-1275, 2007.

or death among 796 men with unstable angina or non-Q-wave myocardial infarction who were treated with low-dose aspirin. 16

The benefits of early aspirin therapy after an ischemic stroke were elucidated in two contemporary large, randomized trials of patients with acute stroke: the Chinese Acute Stroke Trial (CAST) ¹⁷ and the Ischemic Stroke Trial (IST). ¹⁸ In more than 20,000 patients enrolled in CAST, 160 mg of aspirin given within 48 hours of an ischemic stroke prevented 6.8 deaths or recurrent nonfatal strokes per 1000 patients treated. ¹⁷ In a 2 x 2 factorial open-label design, IST investigators examined the effects of subcutaneous heparin, 300 mg of aspirin, or both administered within 48 hours of an ischemic stroke. Aspirin was associated with 11 fewer deaths or recurrent strokes per 1000 patients treated. ¹⁸ The results of these trials, analyzed together, revealed that this benefit was offset slightly by an excess of 2 cases of intracranial hemorrhage per 1000 patients treated. ¹⁹

Treatment with aspirin has also been an essential adjunct in patients undergoing coronary revascularization. Among patients undergoing coronary artery bypass grafting (CABG), aspirin administration both before and soon after surgery has been demonstrated to improve both early and 1-year patency of the saphenous vein graft. ^{20,21} Aspirin administered after coronary angioplasty has been associated with a decreased risk of the composite endpoint of death, restenosis, or myocardial infarction in comparison with placebo (30% versus 41%, respectively). ²² As might be expected, the addition of aspirin to thrombolytic therapy also reduces rates of recurrent ischemia and infarct-related reocclusion of arteries. ²³

The ATC provided irrefutable evidence in favor of aspirin for secondary prevention with a more recent meta-analysis of 195 trials that included more than 135,000 patients. ²⁴ This meta-analysis revealed similar risk reduction with antiplatelet therapy among patients at high risk, and this reduction also extended to patients with stable angina, atrial fibrillation , and peripheral artery disease. ²⁴

Primary Prevention

To date, six large, randomized trials have been undertaken to study aspirin for the primary prevention of cardiovascular disease (Table 7-1). The British Doctors' Trial was conducted to evaluate the effect of a daily 500-mg dose of aspirin in healthy male physicians. ²⁵ Of the 5139 subjects studied, a majority of participants were older than 60 years and were either current or ex-smokers. After 6 years of follow-up, there were no statistically significant differences in the rates of fatal or nonfatal myocardial infarction, stroke, or all-cause mortality between patients assigned to receive aspirin therapy and those assigned to receive no aspirin. There was, however, an approximate 50% reduction in TIA among physicians treated with aspirin. This trial was not

blinded and did not have a placebo control group. Over the course of the study, 44.3% of physicians assigned to receive aspirin therapy discontinued the drug. ²⁵

The Physicians' Health Study, a larger trial designed to assess the efficacy of aspirin in reducing cardiovascular events, enrolled 22,071 male physicians in the United States. ²⁶ In a double-blind, placebo-controlled design, healthy physicians were randomly assigned to receive 325 mg of aspirin every other day or betacarotene in a 2 x 2 factorial design. The study, which was terminated early, demonstrated a significant (44%) risk reduction in the rate of total myocardial infarction. Similarly, there was an 18% risk reduction in the composite outcome of nonfatal myocardial infarction, nonfatal stroke and cardiovascular death. Despite this robust finding, aspirin therapy did not confer a cardiovascular mortality benefit in this study. ²⁶

The effect of aspirin and warfarin in reducing cardiovascular events in 5085 men was evaluated in the Thrombosis Prevention Trial, a randomized, double-blind, placebo-controlled trial. ²⁷ In a 2 x 2 factorial design, men who did not have established cardiovascular disease but were deemed to be at high risk for vascular disease were randomly assigned to receive treatment with aspirin, 75 mg daily, and warfarin with a target international normalized ratio (INR) of 1.5. Aspirin therapy, either alone or in combination with warfarin , conferred a 20% reduction in the primary endpoint of cardiovascular death and fatal and nonfatal myocardial infarction. This reduction was driven primarily by a 32% reduction in the risk of non-fatal myocardial infarction. In a result consistent with those of the large studies preceding the Thrombosis Prevention Trial, no mortality benefit with aspirin therapy was demonstrated. ²⁷

In the Hypertension Optimal Treatment study, the effect of low-dose aspirin was investigated in an international cohort of 18,790 men and women, aged 50 to 80, with hypertension. ²⁸ A separate arm of the study was concerned with the effect of antihypertensive therapy directed at diastolic blood pressure and cardiovascular outcomes. In this randomized, placebo-controlled, double-blind study, a daily dose of 75 mg of aspirin was associated with a 15% relative risk reduction in major cardiovascular events, defined as both fatal and nonfatal stroke, fatal and nonfatal myocardial infarction, or cardiovascular death.

Another trial designed to examine the efficacy of aspirin among men and women at risk for cardiovascular disease

72 was the Primary Prevention Project. ²⁹ In an open-label 2 x 2 factorial design, 4495 patients were randomly assigned to receive a 100-mg daily dose of aspirin, as well as vitamin E. Patients were eligible for inclusion in the trial if they had a history of hypertension, hypercholesterolemia, diabetes, family history of premature coronary artery disease, were obese, or were older than 65. A majority of patients included I in the study had at least two or more of these risks factors. The mean age of participants was more than 60 years, and 57.7% were women. The trial was terminated prematurely

because data from the Hypertension Optimal Treatment and Thrombosis Prevention Trial provided evidence in favor of aspirin for primary prevention. After a mean follow-up period of 3.6 years, investigators demonstrated a 44% relative risk treatment with aspirin. Similarly, they demonstrated a 22%

infarction, or stroke. 30

A large meta-analysis 31 of the six aforementioned primary risk of cardiovascular death.

In an update, the ATC performed another meta-analysis 32 of the six major primary prevention trials, which was strengthened by the availability of individual participant data. In primary prevention studies, aspirin was associated with a 12% reduction in the risk ratio of a major adverse cardiovascular event in comparison with no aspirin (0.51% effect was similar in both men and women and was independent of age. Moreover, the small absolute benefit was counterbalanced by an increase in major extracranial hemorrhage in comparison to no aspirin (0.10% versus 0.07% conferred an absolute reduction of 1.5% per year in the rate of occurrence of a serious vascular event. 32

Although previous primary prevention trials of aspirin 24 included analyzes of data from subgroups of patients with diabetes, these provided insufficient information with regard to efficacy in this important patient population. In the Japanese controlled type 2 diabetes, with a mean age of 65 and without known cardiovascular disease, were randomly assigned to receive 81 or 100 mg daily of aspirin versus no aspirin. 33 The overall event rate was low in this population, and a significant reduction in major cardiovascular events was not observed in the subjects taking aspirin. However, a reduction in the secondary composite endpoint of fatal stroke or myocardial diabetic patients older than 65, a benefit was derived from lowprimary endpoint (6.3% versus 9.2% respectively). 33

In the Prevention of Progression of Arterial Disease and Diabetes (POPADAD) trial, which was contemporaneous with the JPAD trial, the efficacy of low-dose (100 mg) daily aspirin and antioxidant therapy in preventing cardiovascular events was evaluated in 1276 patients with diabetes and asymptomatic peripheral artery disease. ³⁴ The investigators did not observe a statistically significant benefit of aspirin over placebo in this randomized, controlled, double-blind study. In another trial, Aspirin for Asymptomatic Atherosclerosis, 3350 healthy men and women with asymptomatic peripheral artery disease (as defined by an ankle-brachial index < 0.95) were randomly assigned to receive aspirin, 100 mg daily, or placebo. After a mean follow-up period of 8.2 years, no difference in major cardiovascular events or all-cause mortality was observed between participants randomly assigned to receive aspirin or placebo. 35

Several ongoing trials have been designed to address remaining reduction in cardiovascular death among patients assigned to questions regarding the efficacy of aspirin in primary prevention. The plan of A Study of Cardiovascular Events in Diabetes relative risk reduction in the primary endpoint of (ASCEND) is to assess the effectiveness of low-dose daily aspirin in cardiovascular death, nonfatal myocardial infarction, or stroke. 10,000 patients with diabetes. ³⁶ Similarly, the Aspirin and Simvastatin Combination for Cardiovascular Events Prevention The Women's Health Study was designed to address the role Trial in Diabetes (ACCEPT-D) is designed to assess the efficacy of of aspirin in the primary prevention of cardiovascular disease open-label daily aspirin, either alone or in combination with in women. A total of 39,876 female health professionals were simvastatin, in reducing cardiovascular events in approximately randomly assigned to receive 100 mg of aspirin every other 5000 patients with diabetes. ³⁷ Because researchers in most primary day. 30 After a mean follow-up period of 10 years, this treatment prevention studies have examined the benefit of aspirin in was found to confer 17% and 22% reductions in the risk of populations at relatively low risk, the ongoing Aspirin to Reduce stroke and TIAs, respectively, but no significant reduction in Risk of Vascular Events (ARRIVE) trial will address the role of the risk of myocardial infarction or cardio vascular death. In the aspirin in an international cohort of approximately 12,000 patients subgroup of women aged 65 and older, however, aspirin was deemed to be at moderate risk (20% to 30%) of developing a associated with a significant (26%) reduction in the risk of the cardiovascular event over 10 years. 38 The role of aspirin in the primary endpoint of cardiovascular death, nonfatal myocardial primary prevention of cardiovascular disease in elderly patients is being studied in the Aspirin in Reducing Events in the Elderly (ASPREE) trial. 39 The aim of the Japanese Primary Prevention prevention trials concluded that aspirin reduces composite Project with Aspirin is to evaluate cardiovascular outcomes in 10,000 cardiovascular events by 12% and 14% in women and men, Japanese patients older than 60 with at least one additional respectively. Overall, aspirin was not associated with a lower traditional cardio vascular risk factor who are treated with 100 mg of aspirin daily. 40

Dosage

Daily aspirin doses of only 30 mg have been demonstrated to completely inhibit synthesis of platelet thromboxane. 41 Despite this observation, the optimal dose of aspirin for an individual patient is versus 0.57% per year, respectively). The magnitude of this not known. Some investigators have speculated that higher dosages of aspirin may paradoxically attenuate the antithrombotic effect of thromboxane inhibition by causing inhibition of the vasodilator prostacyclin. 42 However, a wide variety of aspirin dosages (ranging from 50 to 1500 mg) have been demonstrated to be efficacious for per year, respectively). For secondary prevention, aspirin prevention of cardiovascular events, and a formal comparison of several different doses has never been performed in the context of a randomized, controlled, prospective trial in coronary artery disease.

Results of one ATC meta-analysis suggested similar reduction in vascular events across a wide range of aspirin dosages. 24 In a post hoc analysis of the Clopidogrel in Unstable Angina to prevent Primary Prevention of Atherosclerosis with Aspirin for Recurrent Events (CURE) study, increasing dosages of aspirin (< 100 Diabetes (JPAD) trial, 2539 men and women with well- mg, 101 to 199 mg, and > 200 mg daily) administered with either placebo or clopidogrel were not associated with greater clinical benefit. 43 Moreover, higher rates of major bleeding were observed with escalating dosages of aspirin compared with placebo (1.9%, 2.8%, and 3.7%, respectively). ⁴³ A meta-analysis of 31 randomized trials that included more than 192,000 patients reached a similar conclusion; the risk of major bleeding events was lowest in patients who took the lowest aspirin dosage. 44 This finding was confirmed in infarction was observed. In a prespecified subgroup analysis of an observational study of participants in the Clopidogrel for High Atherothrombotic Risk and Ischemic Stabilization (CHARISMA) dose aspirin in comparison with the control condition in the trial. 45 In this large prospective study of patients at high risk of cardiovascular events, patients

the composite outcome of myocardial infarction, death, or stroke expected to heighten the risk of gastrointestinal hemorrhage. 56 with increasing doses of aspirin. In fact, there was a suggestion of significant. 45

mg daily for 6 days and sub sequent maintenance dosing of 75 mg 75 mg daily plus esomepra zole placebo twice daily versus 80 mg infarction, or stroke at 30 days in patients undergoing percutaneous coronary intervention (PCI). 46 No difference in efficacy or hemorrhagic risk was observed between patients who received lowdose aspirin and those who received high-dose aspirin. 46 A prostacyclin among patients who experience adverse cardiac events and those who do not. 47

Formulations

Aspirin exists in a "regular" form, as well as in buffered and entericcoated preparations. Aspirin is rapidly absorbed in the stomach and small intestine after ingestion; inhibition of portal platelet COX enzyme occurs before complete systemic absorption. 48 Levels of aspirin in the systemic circulation peak within 40 minutes after ingestion of regular aspirin and 3 to 4 hours after ingestion of an gastrointestinal complications. 61 enteric-coated preparation. 49 A pharmacodynamic study in 12 healthy volunteers demonstrated that near-maximal platelet Drug Interactions thromboxane inhibition, occurring over a mean of 13.6 minutes, is achieved most efficiently when a 325-mg aspirin tablet is chewed. 50 Swallowing a whole buffered tablet doubles the time necessary to achieve maximal platelet inhibition. 50 Although enteric-coated preparations may have theoretical benefit in reducing gastric irritation and bleeding, the risk of gastrointestinal bleeding observed with aspirin is also increased because of its systemic effect. Entericcoated aspirin does not seem to confer protection against aspirin and ibuprofen. The latter combination was associated with gastrointestinal bleeding in comparison with buffered or regular preparations of the same dose. 51

Hemorrhagic Complications

The most feared complications of antiplatelet therapy are sequelae from hemorrhage. The majority of bleeding complications arise from the gastrointestinal tract; the estimated relative risk was 2.1 in one meta-analysis of 22 randomized primary or secondary prevention studies in which 75- to 325-mg dosages of aspirin were compared with placebo. 52 The relative risk of intracranial hemorrhage was 1.7; no differ ences between major bleeding and dosages of aspirin were observed. This translated to an annual absolute increase in major bleeding of 0.12%. 52 In a prospective, observational study of 991 patients with coronary artery disease who were treated with 75- to 300-mg of aspirin, the incidence of upper gastrointestinal hemorrhage was 1.5% over 2 years of follow-up. 53 It has been estimated that aspirin contributes to an excess of 5 cases of gastrointestinal hemorrhage per 1000 patients treated. 54 Gastric toxicity, as measured by inhibition of gastric prostaglandin

were randomly assigned to receive clopidogrel or placebo in synthesis, is thought to be dose dependent, and a 50% reduction in addition to background aspirin therapy (at daily doses of 162 mg or gastric prostaglandin is observed at dosages as low as 30 mg/day. 55 lower). A post hoc analysis demonstrated no significant reduction in Therefore, all dosages currently prescribed in clinical practice can be

The extent to which this risk can be attenuated by proton pump harm to patients treated with higher doses of aspirin in addition to inhibition has been examined in both asymptomatic patients and clopidogrel, with increased rates of cardiovascular events and a those with prior gastroduodenal ulcers. In a prospective, doublegreater incidence of bleeding, although this was not statistically blind study of more than 900 asymptomatic patients requiring lowdose aspirin therapy, use of a proton pump inhibitor (PPI) for 26 Results of the Clopidogrel Optimal Loading Dose Usage to weeks was associated with a lower rate of endoscopic ulcers than Reduce Recurrent Events/Optimal Antiplatelet Strategy for was placebo (1.6% versus 5.4%, respectively). ⁵⁷ In another study of Interventions (CURRENT-OASIS 7) trial have enhanced the 123 patients with recently healed gastroduodenal ulcers and treated understanding of both aspirin and clopidogrel dosing in acute *Helicobacter pylori* infection, the combination of 100 mg of aspirin coronary syndromes (ACS). 46 In a double-blind 2 x 2 factorial design, and 30 mg of lansoprazole was associated with fewer recurrent approximately 25,000 patients with ACS treated with an early ulcer complications was the combination of aspirin and placebo invasive strategy were randomly assigned to receive conventional over 1 year (1.6% versus 14.8%, respectively). 58 In a similar clopidogrel dosages (300-mg loading dose, fol lowed by 75 mg daily) randomized, placebo-controlled trial of 320 patients with a recent versus high-dose clopidogrel (600-mg loading dose, followed by 150 bleeding ulcer, investigators studied the combination of clopidogrel daily). 47 Subjects in each of these groups were further randomly aspirin plus esomepra zole 20 mg twice daily. 59 Clopidogrel was assigned, in an open-label manner, to receive high-dose aspirin (300 associated with a higher rate of recurrent bleeding over 1 year than to 325 mg) or low-dose aspirin (75 to 100 mg) after an initial 300-mg was the combination of aspirin and PPI (0.7% versus 8.6%, dose of aspirin. High-dose clopidogrel was associated with a respectively). 59 During 12 weeks of follow-up, the histamine H2 significant reduction in the composite of death, myocardial receptor antagonist famotidine was demonstrated to decrease the risk of endoscopic esophagitis and peptic ulcers in comparison with placebo in patients who received 75 to 325 mg of daily aspirin

By consensus, the American College of Cardiology Foundation prespecified substudy will also be conducted to examine the effects (ACCF), American College of Gastroenterology (ACG), and of aspirin dosing on urinary metabolites of thromboxane and American Heart Association (AHA) recommend reducing chronic aspirin dosages to 81 mg daily with the addition of a daily dose of a PPI in patients with a history of gastrointestinal hemorrhage or ulcer or in patients at risk of these complications, such as those who take maintenance steroid medication, elderly patients, or patients with a history of dyspepsia. 61

In addition, testing and treatment of H. pylori is advocated initiation of chronic antiplatelet therapy in patients with a history of peptic ulcer disease. Replacing aspirin with clopidogrel is not recommended as a strategy for reducing the risk of recurrent

Other nonsteroidal anti-inflammatory drugs (NSAIDs) can interact in deleterious ways with aspirin. The addition of NSAIDs to aspirin potentiates the risk of gastrointestinal events. However, concomitant NSAID use may also mitigate the protective effect of aspirin. MacDonald and Wei 62 reported on trends in mortality among more than 7000 patients with cardiovascular disease discharged from the hospital with prescriptions for aspirin or for the combination of an excess risk of both all-cause mortality and cardiovascular death (hazard ratios, 1.9 and 1.7, respectively). The potential mechanism of this interaction was evaluated in healthy volunteers who were administered ibuprofen, followed by 81 mg of aspirin. 63 Ibuprofen administered before aspirin or several times daily blocked normal aspirin-induced platelet inhibition. However, the administration of aspirin 2 hours before a single dose of ibuprofen resulted in expected irreversible COX-1 inhibition. 63 Naproxen has also been demonstrated

FIGURE 7-2 Possible mechanisms of aspirin resistance. COX, cyclooxygenase; GP, glycoprotein; mRNA, messenger ribonucleic acid; PGF_{2a}, prostaglandin factor 2a; vWF, von Willebrand factor. (*From Bhatt DL: Aspirin resistance: more than just a laboratory curiosity*, J Am Coll Cardiol 43:1127-1129, 2004.) to antagonize the COX-1 inhibition of aspirin in vitro, presumably by functioning as a competitive inhibitor of the COX enzyme. ⁶⁴ Amplifying concerns about NSAIDs as a class, a large Finnish case-control study demonstrated a significant increase in the risk of first myocardial infarction with use of either conventional or selective COX-2 inhibitor NSAIDs. ⁶⁵ . ⁶⁶

Aspirin Resistance

Despite appropriate doses of aspirin, many patients develop recurrent ischemic events. This clinical dilemma has often been attributed to aspirin resistance, a broad term that encompasses the wide variety of factors thought to contribute to this phenomenon (Figure 7-2). At the simplest level, patients' nonadherence to aspirin therapy, underprescription by physicians, drug interaction with ibuprofen or naproxen, and malabsorption may all play a role. 67 It is also known that platelet activation can occur via thromboxane-independent pathways. 68 One such mechanism may involve COX independent production of the arachidonic acid derivative 8-iso-prosta-glandin factor F 2 , PGF _{2a} a potent vasoconstrictor and platelet aggregant, released in response to oxidative stress. 69 Because aspirin is a relatively weak inhibitor of COX-2, it has also been postulated that platelet COX-2, normally expressed in response to inflammatory stimuli, may result in sufficient synthesis of thromboxane A 2 to contribute to aspirin resistance. 70 Other genetic factors may also contribute to observed differences in platelet responsiveness. The platelet polymorphism PI A2 has been associated with aspirin resistance. 68 Aspirin resistance has been observed in patients with acute myocardial infarction 71 and elicited by exercise in patients with stable coronary artery disease. 72 One systematic review of 15 studies revealed a wide range in estimates of the prevalence of laboratory aspirin resistance (5% to 65%). 73 The lack of a uniform definition of aspirin resistance and its measurement has limited the understanding of this entity. The "gold standard" test of platelet function, light transmission aggregometry, is the most precise; however, it is time consuming and cannot be performed at the patient's bedside. 74

The implications of inadequate aspirin-induced platelet inhibition were assessed in a nested case-control study of participants in the Heart Outcomes Prevention Evaluation (HOPE). Eikelboom and colleagues 75 found an independent

association between increasing urinary thromboxane levels, a marker of aspirin resistance, and major cardiovascular events. In another prospective study of 326 patients with stable cardiovascular disease, aspirin resistance, as measured by a onetime optical platelet aggregation test, was present in 5.2% of patients and associated with a significant increase in the rate of the combined endpoint of myocardial infarction, stroke, or death in comparison with patients not deemed resistant (24% versus 10%, respectively). ⁷⁶ In data congruent with these findings, Chen and colleagues 77 demonstrated an almost threefold increase in the risk of periprocedural myocardial infarction in patients undergoing nonurgent PCI who were deemed aspirin resistant according to a commercial point-of-care assay. More recently, a prespecified analysis of the CHARISMA trial confirmed the findings of the HOPE substudy and revealed an increased risk of stroke, myocardial infarction, or death in patients whose urinary 11-dehydro-thromboxane B 2 levels were in the highest quartile. ⁷⁸ More over, clopidogrel (in subjects who received it) did not appear to attenuate this relationship. Interestingly, female sex, increasing age, peripheral artery disease, tobacco use, and use of angiotensin-converting enzyme inhibitors or oral hypoglycemic agents were independently associated with incomplete thromboxane inhibition. 78

Guidelines

The US Preventive Services Task Force (USPSTF) recommends the use of aspirin in men aged 45 to 79 and women aged 55 to 79 for the primary prevention of a cardiovascular event if the perceived benefit of aspirin outweighs the potential harm caused by an increased risk of gastrointestinal haemorrhage (Figure 7-3). ⁷⁹ For patients at moderate risk for cardiovascular events, the American College of Chest Physicians (ACCP) recommends 75 to 100 mg of aspirin daily. ⁸⁰ The American Diabetes Association advocates the use of 75 to 162 mg of aspirin daily in patients with diabetes who are older than 40 or for those who have other traditional risk factors for cardiovascular disease. ⁸¹ For patients with peripheral artery disease, the American College of Cardiology (ACC)/AHA recommends 75 to 325 mg of aspirin daily for the prevention of stroke, myocardial infarction, or cardiovascular death. ⁸²

After an ST-elevation myocardial infarction (STEMI), the ACCP recommends initiation of aspirin at a dose of 160 to 325 mg, with subsequent reduction to 75 to 100 mg, to be continued indefinitely. ⁸⁰ The ACCP also recommends indefinite low-dose aspirin (75 to 100 mg daily) after PCI, CABG, carotid endarterectomy, and peripheral revascularization. ⁸⁰ A focused update of the 2004 STEMI guidelines recommends aspirin initiation at a dose of 162 to 325 mg for 1, 3, and 6 months after bare-metal, sirolimus, and paclitaxel drug-eluting stents, respectively, with reduction to 75 to 162 mg daily thereafter (ACC/AHA Class 1, level of evidence B). ⁸³ The AHA/ American Stroke Association (ASA) recommends administration of 325 mg of aspirin within 24 to 48 hours of an acute ischemic stroke, except for patients receiving thrombolytic therapy, for whom aspirin should be deferred 24 hours. ⁸⁴

THIENOPYRIDINES

Mechanism of Action

The thienopyridines, of which ticlopidine and clopidogrel are the prototypes, irreversibly inhibit platelets by binding to P2Y 12, the G protein-coupled receptor that is normally activated by ADP released from injured endothelium and red blood cells. 85 Through interaction with the P2Y 12 and P2Y 1 platelet receptors, ADP triggers a cascade of events that result in platelet aggregation and in further release of ADP from the activated platelet, thus potentiating the initial response. 86,87

ASPIRIN FOR THE PREVENTION OF CARDIOVASCULAR DISEASES CLINICAL SUMMARY OF US PREVENTIVE SERVICES TASK FORCE RECOMMENDATIONS

Population	Men Age 45-79 Years	Women Age 55-79 Years	Men Age <45 Years	Women Age <55 Years	Men and Women Age >80 Years
Recommendation	Encourage aspirin use when potential CVD benefit (MIs prevented) outweighs potential harm of GI hemorrhage	Encourage aspirin use when potential CVD benefit (strokes prevented) outweighs potential harm of GI hemorrhage	Do not encourage aspirin use for MI prevention	Do not encourage aspirin use for stroke prevention	recommendation
	Grade:	A	Gra	de: D	Grade: I (insufficient evidence)

Shared decision making is strongly encouraged with individuals whose risk is close to (either above or below) the estimates of 10-year risk levels indicated below. As the potential CVD benefit increases above harms, the recommendation to take aspirin should become stronger.

To determine whether the potential benefit of MI's prevented (men) and strokes prevented (women) outweighs the potential harm of increased GI hemorrhage, both 10-year CVD risk and age must be considered.

The table above applies to adults who are not taking NSAIDs and who do not have upper GI pain or a history

How to Use This

Risk Level at Which CVD Events Prevented (Benefit) Exceeds GI Harm					
ı	Men	Women			
Age	10-Year CHD Risk	Age	10-Year Stroke Risk		
45-59 years	>4%	55-59 years	>3%		
60-69 years	>9%	60-69 years	>8%		
70-79 years	>12%	70-79 years	>11%		

NSAID use and history of GI ulcers increase the risk for serious GI bleeding events considerably and should be considered in determining the balance of benefits and harm.

NSAID use combined with aspirin use approximately quadruples the risk for serious GI bleeding events compared with the risk with aspirin use alone. The rate of serious bleeding in aspirin users is approximately 2 to 3 times greater in patients with a history of GI ulcers

For men: Risk factors for CHD include age, diabetes, total cholesterol level, HDL cholesterol level, blood pressure, and smoking.

CHD risk estimation tool: http://healthlink.mcw.edu/article/923521437.html

Risk Assessment

For women: Risk factors for ischemic stroke include age, high blood pressure, diabetes, smoking, history of CVD, atrial fibrillation, and left ventricular hypertrophy.

Stroke risk estimation tool: www.silentstroke.org/PersonalStrokeRisk1.xls

Relevant Recommendations from USP-STF The USPSTF has made recommendations on screening for abdominal aortic aneurysm, carotid artery stenosis, CHD, high blood pressure, lipid disorders, and peripheral arterial disease. These recommendations are available at www.preventiveservices.ahrg.gov.

FIGURE 7-3 US Preventive Services Task Force (USPSTF) recommendations for aspirin use in primary prevention. CHD, coronary heart disease; CVD, cardiovascular disease; GI, gastrointestinal; HDL, high-density lipoprotein; MI, myocardial infarction; NSAID, nonsteroidal anti-inflammatory drug. (From US Preventive Services Task Force: Aspirin for the prevention of cardiovascular disease: clinical summary of US Preventive Services Task Force Recommendation, AHRQ Publication No. 09-05129-EF-3, Rockville, MD, March 2009, Agency for Healthcare Research and Quality. Available at: http://www.ahrq.gov/clinic/uspstf09/aspirincvd/ aspcvdsum.htm.)

Ticlopidine

Ticlopidine, first studied in humans in 1975, inhibits ADP-induced platelet aggregation in a dose-dependent manner, with an onset of action of 24 to 48 hours. ⁸⁸ Like clopidogrel, ticlopidine is a prodrug and must be metabolized by the cytochrome P-450 system to an active metabolite. ⁸⁹ Although many early trials provided evidence to support the use of ticlopidine in patients with established cardiovascular disease, adverse hematological side effects and rather slow onset of action in comparison with clopidogrel have curtailed its widespread subsequent use. Among patients taking ticlopidine, serious neutropenia has been reported in fewer than 1% to as high as 3.4%, ⁹⁰⁻⁹⁴ and thrombotic thrombocytopenic purpura has been reported in 0.02%. ⁹⁵

Two such early trials were the Canadian American Ticlopidine Study (CATS) 90 and the Ticlopidine Aspirin Stroke

Study (TASS). ⁹² In the CATS trial, more than 1000 patients with recent thromboembolic stroke were randomly assigned to receive treatment with ticlopidine or placebo; a nearly 25% relative risk reduction was demonstrated in the rate of the combined endpoint of vascular death, myocardial infarction, or death. ⁹⁰ In TASS, ticlopidine was compared with high-dose aspirin in more than 3000 patients who had suffered a recent neurological event; a 21% relative risk reduction was demonstrated in the rate of recurrent fatal and

nonfatal stroke in favor of ticlopidine. ⁹² Incongruent with this result were the findings of the African-American Antiplatelet Stroke Prevention Study (AAASPS), which did not show a reduction in the composite endpoint of stroke, myocardial infarction, or vascular death in African American patients treated with ticlopidine after an ischemic stroke. ⁹¹

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The Swedish Ticlopidine Multicentre Study demonstrated a 29% reduction in all-cause mortality among 687 patients with established peripheral artery disease treated with ticlopidine, in comparison with placebo. 93 This mortality benefit was explained entirely by a reduction in fatal myocardial infarction. The early use of ticlopidine was further supported by a study that demonstrated a nearly 47% relative risk reduction in vascular death among 653 patients with unstable angina treated with ticlopidine in an open-label trial. 96 The additional antiplatelet benefit of ticlopidine was later demonstrated to extend to PCIs, previously complicated by stent thrombosis in the era of single antiplatelet therapy and oral anticoagulation. 94,97,98 Soon after, the results of the Clopido grel Aspirin Stent International Cooperative Study (CLAS SICS) suggested superiority of the combination of aspirin and clopidogrel over that of aspirin and ticlopidine in patients undergoing placement of coronary stents. ⁹⁹ Although this study was not statistically powered to compare the efficacy of these two antiplatelet regimens, the combination of aspirin and clopidogrel was associated with significantly fewer non cardiac adverse effects than was the combination of ticlopidine and aspirin (4.6% versus 9.1%, respectively). 99 More conclusive evidence arose from a meta-analysis of both registry and randomized trial data in which clopidogrel and ticlopidine were compared: The rate of major adverse cardiac events was reduced 50% with combination clopidogrel and aspirin in comparison with ticlopidine and aspirin. 100

Clopidogrel

Secondary Prevention

Clopidogrel has been tested in the secondary prevention of cardiovascular disease in several trials (Table 7-2). The -Clopidogrel versus Aspirin in Patients at Risk of Ischemic Events (CAPRIE) study was the first large, randomized, placebocontrolled trial to test the efficacy of clopidogrel in preventing cardiovascular events. 101 This international, multicenter study included 19,185 patients, predominantly male, with a mean age of 63 who had sustained a recent myocardial infarction, stroke, or symptomatic peripheral artery disease. The subjects were monitored for a mean of almost 2 years. Clopidogrel (75 mg daily) conferred an 8.7% relative risk reduction in the rate of the composite endpoint of myocardial infarction, stroke, or vascular death in comparison with a daily 325-mg dose of aspirin. 101 In subgroup analyzes of patients with diabetes and prior CABG in the CAPRIE trial, clopidogrel was also more effective than aspirin in reducing the rate of the combined endpoint of vascular death, myocardial infarction, or stroke. 102,103

The salutary effect of dual-antiplatelet therapy with clopidogrel and aspirin in patients with ACS was established in the CURE trial. 104 Among 12,562 men and women with unstable angina or non-ST-elevation myocardial infarction, a 300-mg loading dose of clopidogrel followed by a dose of 75 mg of clopidogrel daily with open-label aspirin therapy (75 to 325 mg) was associated with a lower rate of the combined endpoint of cardiovascular death, myocardial infarction, or stroke than was placebo (9.6% versus 11.4%, respectively). The protective effect of clopidogrel was evident within the first 24 hours after randomization, and clopidogrel also reduced the risk of inhospital ischemia, recurrent angina, revascularization, and heart failure. 104 The additional early benefit of clopidogrel in comparison with placebo was also shown to extend to patients in the CURE study who subsequently underwent CABG 105 and PCI. 106 Another CURE substudy demonstrated the consistent benefit of clopidogrel across various risk groups, as defined by the Thrombolysis In Myocardial Infarction (TIMI) risk score. 107

The use of dual-antiplatelet therapy with clopidogrel and aspirin before PCI was supported by evidence from the Clopidogrel for the Reduction of Events During Observation (CREDO) trial. ¹⁰⁸ In 2116 patients undergoing elective PCI, pretreatment with a 300-mg loading dose of clopidogrel followed by 1 year of dual-antiplatelet therapy (clopidogrel 75 mg daily and aspirin 81-325 mg daily) was associated with a nearly 27% relative risk reduction in the composite end point of myocardial infarction, death, or need for target vessel revascularization in comparison with placebo. ¹⁰⁸

On the basis of the premise that combination therapy with clopidogrel and aspirin might attenuate cardiovascular risk beyond that observed with clopidogrel alone, the CHA RISMA trial was conducted to evaluate the efficacy of clopidogrel and low-dose aspirin for the prevention of major cardiovascular events. ¹⁰⁹ The 15,603 patients with established cardiovascular disease or multiple cardiovascular risk factors were monitored for a median of 28 months. Clopidogrel and aspirin did not result in significant benefit with regard to the composite endpoint of stroke, myocardial infarction, or cardiovascular death in comparison with placebo plus aspirin. ¹⁰⁹ However, in a subsequent analysis of patients with prior myocardial infarction, symptomatic peripheral artery disease, or stroke, the combination of clopidogrel and aspirin afforded a 1.5% absolute risk reduction in the composite endpoint of stroke, myocardial infarction, or cardio vascular death. ¹¹⁰

The additional benefit of clopidogrel among patients with STEMI has been demonstrated in two large-scale randomized, placebo-controlled trials. 111,112 The Clopidogrel and Metopro lol in Myocardial Infarction Trial (COMMIT) enrolled 45,852 Chinese patients with an acute myocardial infarction to receive clopidogrel, 75 mg daily, or placebo in addition to aspirin, 162 mg daily. 111 Also in this 2 x 2 factorial design, the effect of metoprolol (intravenous followed by oral preparations) was evaluated. Clopidogrel treatment for a mean of approximately 2 weeks was associated with a 9% odds reduction in the composite of myocardial infarction, stroke, or death, as well as a 7% odds reduction in all-cause mortality. 111 Similarly, the Clopidogrel as Adjunctive Reperfusion Therapy-Thrombolysis In Myocardial Infarction 28 (CLARITY-TIMI 28) study demonstrated a 36% odds reduction in the composite endpoint of death, myocardial infarction, or infarct-related artery occlusion, demonstrated angiographically, in 3491 patients with STEMI who were treated with clopidogrel and fibrinolytics. 112 This reduction was achieved without a significant increase in the risk of major bleeding in both trials. In a prespecified analysis of patients who underwent PCI in the CLARITY-TIMI 28 study, random assignment to pretreatment with a 300-mg loading dose of clopidogrel was associated with a 46% odds reduction at 30 days in the composite endpoint of stroke, myocardial infarction, or death. 113

The role of clopidogrel in the prevention of cerebrovascular events has been addressed in a prospective manner. Diener and colleagues ¹¹⁴ studied the addition of low-dose aspirin to background clopidogrel therapy among patients with recent stroke or TIA and at least one additional cardiovascular risk factor in the Management of Atherothrombosis with Clopidogrel in High-Risk Patients (MATCH) study. After 18 months of treatment, there was no statistically significant benefit of dual-antiplatelet therapy with regard to stroke, myocardial infarction, or vascular death, and an important increase in major bleeding was observed in comparison with clopidogrel and placebo. ¹¹⁴ With this background, the Prevention Regime

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	No. of					Relative Risk	
rial .	Patients 19,185	Population Patients with established vascular disease	Follow-Up Average: 1.9 years		Primary Endpoint Stroke, vascular death, or MI	Reduction	P 0.043
Clopidogrel vs. Aspirin in Patients at Risk of Ischemic Events (CAPRIE)				·			
Clopidogrel in Unstable Angina to prevent Recurrent Events (CURE)	12,562	Patients with ACS	3-12 months	Clopidogrel + aspirin versus aspirin alone	CV death, nonfatal MI, or stroke	20%	< 0.00
PCI-CURE	2658	Patients with NSTEMI and undergoing PCI	30 days	Clopidogrel + aspirin versus aspirin alone	CV death, MI, urgent TLR	30%	0.03
Clopidogrel and Metoprolol in Myocardial Infarction Trial (COMMIT)	45,852	Patients with acute MI	Mean: 15 days	Clopidogrel + aspirin versus aspirin alone	Death, stroke, or reinfarction	9%	0.002
Clopidogrel as Adjunctive Reperfusion Therapy- Thrombolysis In Myocardial Infarction (CLARITY-TIMI 28)	3491	Patients with STEMI and receiving fibrinolytics	Angiography after a median of 84 hours	Clopidogrel + aspirin versus aspirin alone	Death, MI, or heart attack- related occlusion of artery	31%	< 0.00
Clopidogrel for the Reduction of Events During Observation (CREDO)	2116	Patients undergoing elective PCI or with high likelihood of needing PCI	1 year	Clopidogrel + aspirin versus aspirin alone	Death, stroke, or MI	26.9%	0.02
CI CLARITY	1863	Patients with STEMI and receiving fibrinolytics and PCI	30 days after PCI	Clopidogrel + aspirin versus aspirin alone	Death, recurrent MI, or stroke	42%	0.008
Clopidogrel for High Atherothrombotic Risk and Ischemic Stabilization (CHARISMA)	15,603	Patients with known CV disease or multiple risk factors for CV disease	Median: 28 months	Clopidogrel + aspirin versus aspirin alone	CV death, stroke, or MI	7%	0.22
Management of Atherothrombosis with Clopidogrel in High-Risk Patients (MATCH	7599	Patients with recent TIA or stroke and risk factor for stroke	18 months	Clopidogrel + aspirin versus clopidogrel alone	Ischemic stroke, vascular death, MI, or rehospitalization for ischemia	6.4%	0.244
Atrial Fibrillation Clopidogrel Trial with Irbesartan for Prevention of Vascular Events (ACTIVE)- W	6706	Patients with AF and risk factors for stroke	Median: 1.28 years	Clopidogrel + aspirin versus warfarin	Stroke, MI, vascular death, or non-CNS systemic embolus	- 44%	< 0.00
ACTIVE-A	7554	Patients with AF and risk factors for stroke but ineligible for VKA	Median: 3.6 years	Clopidogrel + aspirin versus aspirin alone	Stroke, MI, vascular death, or non-CNS systemic embolus	11%	0.01

ACS, acute coronary syndrome; AF, atrial fibrillation; CNS, central nervous system; CV, cardiovascular; MI, myocardial infarction, NSTEMI, non-ST-elevation myocardial infarction; PCI, percutaneous coronary intervention; STEMI, ST-elevation myocardial infarction; TIA, transient ischemic attack; TLR, target lesion revascularization; VKA, vitamin K antagonists.

Adapted from Meadows T, Bhatt DL: Clinical aspects of platelet inhibitors and thrombus formation, Circ Res 100:1261-1275, 2007.

for Effectively Avoiding Second Strokes (PRoFESS) investigators randomly assigned more than 20,000 men and women with a mean age of 66 who had suffered a recent ischemic stroke to receive either fixed-dose aspirin, 25 mg, and extended-release dipyridamole, 250 mg, twice daily or clopi dogrel, 75 mg daily. 115 The effect of telmisartan was also studied in a 2 x 2 factorial design. After a mean follow-up period of 2.5 years, there was no statistical difference in

either the rate of the primary endpoint of recurrent stroke or the secondary composite endpoint of vascular death, stroke, or myocardial infarction. 115

The first prospective investigation of the role of dual antiplatelet therapy in preventing cardiovascular events in patients with atrial fibrillation was part of the Atrial Fibrilla tion Clopidogrel Trial with Irbesartan for Prevention of Vascular -Events (ACTIVE) family of studies. 116, 117 The ACTIVE-W

78 trial enrolled 6706 patients with atrial fibrillation and additional risk factors for stroke, with a mean CHADS 2 (congestive heart failure, hypertension, age > 75 years, diabetes mellitus, and either stroke or TIA) score of 2, to test the hypothesis that combination clopidogrel and low-dose aspirin would be noninferior to oral anticoagulation with vitamin K antagonists targeted to an INR goal of 2 to 3. 116 The trial was halted prematurely because the combination of clopidogrel and aspirin was associated with an excess risk of the composite endpoint of stroke, myocardial infarction, vascular death, 7 or systemic embolus in comparison with oral anticoagulation (5.6% versus 3.9%, respectively). The superiority of oral anticoagulation was largely driven by a significant reduction in the risk of stroke and systemic embolism. 116 As expected, maintenance of a therapeutic INR is an important is metabolized to inactive metabolites. 128 The remainder must proviso. 118

The superiority of oral anticoagulation over dual antiplatelet therapy in stroke prevention has also been demonstrated in subgroups of ACTIVE-W participants who were at relatively lower risk. 119 Despite the clear superiority of oral anticoagulation over antiplatelet therapy in patients at high risk with atrial fibrillation, it is not appropriate for certain patients. The ACTIVE-A trial, in which 7554 such patients were randomly assigned to receive either clopidogrel or placebo with the background of aspirin, demonstrated a 28% reduction in the risk of stroke with clopidogrel and aspirin. 117 However, dualantiplatelet therapy resulted in a 51% increase in the risk of major extracranial hemorrhage. 117 In the ongoing Secondary Prevention of Small Subcortical Strokes (SPS3) trial, researchers aspirin and clopidogrel in comparison with aspirin and placebo in the prevention of recurrent stroke in patients with lacunar strokes. 120

placement, the optimal duration of this therapy, particularly after drug-eluting stent placement, remains a subject of great debate. The most feared complication after stent placement is stent thrombosis, an uncommon but highly morbid event. An early meta-analysis focused on data from 6675 patients enrolled in randomized trials in which first-generation drug-eluting stents were compared with bare-metal stents; the data revealed a significant increase in the risk of late stent thrombosis in patients treated with drug-eluting stents and less than 6 months of dual-antiplatelet therapy. 121 Although the risk of early stent thrombosis (< 30 days) appeared similar, the risk of stent thrombosis after 1 year was almost five times higher in patients treated with drug-eluting stents. 121 Elaborating on these findings, a registry analysis of more than 4000 patients receiving drug-eluting or bare-metal stents demonstrated a significantly lower rate of the combined endpoint of death or myocardial infarction in patients with drug-eluting stents who were treated with extended clopidogrel than in those treated with 6 or 12 months of clopidogrel. 122 In another observational study of patients treated with drug-eluting stents, the overall rate of stent thrombosis was 1.9% during 18 months of followup, and the major predictor of stent thrombosis within 6 months of placement of a drug-eluting stent was discontinuation of clopidogrel. 123

The possibility of rebound phenomena after cessation of clopidogrel was raised in a retrospective study of more than 3000 patients treated with clopidogrel after ACS. 124 In this Veterans Affairs cohort of patients treated with treated with either medical therapy or PCI, increased rates of the combined endpoint of all-cause mortality or acute myocardial infarction were observed in both medically treated and post-PCI patients after cessation of clopidogrel. Interestingly, there was a grouping of events in the first 90 days after clopidogrel cessation, which raised the specter of a rebound effect. 124 This possibility was supported by in vitro upregulation of proinflammatory markers and increased platelet aggregation after clopidogrel withdrawal in a small study of patients with

diabetes. 125 On the other hand, biological rebound is less likely with an irreversible antiplatelet agent.

Clopidogrel Resistance

An important clinical conundrum arises from the great variability observed in platelet responsiveness to clopidogrel. 126 A growing body of evidence suggests that clopidogrel resistance is associated with poorer cardiovascular outcomes. In a small study of 60 patients with STEMI, hyporesponsiveness to clopidogrel was observed in up to 25% of patients and associated with greater risk of a recurrent cardiovascular event over a 6-month follow-up period. 127 The pharmacogenetic factors underlying this observation have been further elucidated: Approximately 80% of the prodrug clopidogrel undergo hepatic metabolism through a two-step cytochrome P-450dependent process. Among healthy volunteers, Mega and colleagues 129 demonstrated a 30% prevalence of the CYP2C19 allele, a genetic polymorphism that confers loss of function and hence a reduction of the active metabolite of clopidogrel. These investigators also examined the relationship between presence of the CYP2C19 polymorphism and clinical outcomes among 1477 participants assigned to receive clopidogrel in the Trial to Assess Improvement in Therapeutic Outcomes by Optimizing Platelet Inhibition with Prasugrel-Thrombolysis In Myocardial Infarction 38 (TRITON-TIMI 38). In this retrospective analysis, there was a 54% increase in the risk of the composite endpoint of myocardial infarction, car diovascular death, or stroke among carriers of at least one CYP2C19 allele over that of noncarriers. Presence of the CYP2C19 allele was also will examine the efficacy of dual-antiplatelet therapy with 129 These findings were supported by a contemporary report from associated with a threefold increase in the risk of stent thrombosis. the French registry of Acute ST Elevation Myocardial Infarction and Non-ST-Elevation Myocardial Infarction (FAST-MI). ¹³⁰ In patients with an acute myocardial infarction who underwent PCI, the and aspirin has become the standard of care after PCI and stent more than a threefold increase in the risk of adverse cardiovascular events. 130 A genome-wide association study confirmed that this allele may affect clopidogrel response. 131,132

> Despite these data, the optimal management strategy for patients with apparent clopidogrel resistance is not known. In a small case series of 7 patients with stent thrombosis and clopidogrel resistance measured by platelet reactivity, escalation of clopidogrel maintenance doses did not result in improved platelet responsiveness. 128 The ongoing Gauging Responsiveness with a VerifyNow Assay-Impact on Thrombosis and Safety (GRAVITAS) trial is a randomized, placebo-controlled study that should add important information in this regard. 133 With the use of a point-ofcare assay, approximately 2200 patients with high platelet reactivity will be randomly assigned to receive conventional dosages of clopidogrel (75 mg daily) versus 150 mg daily after placement of a drug-eluting stent for 6 months and will be monitored for the occurrence of nonfatal myocardial infarction, cardiovascular death, or stent thrombosis. 133

> Drug-drug interactions may also contribute to clinically observed clopidogrel resistance. Because clopidogrel must be hepatically metabolized through a cytochrome P-450-dependent process, coadministration with CYP450 substrates has also been implicated in reducing the efficacy of clopidogrel. Although results of a small early study of 44 patients undergoing elective stent implantation suggested such an interaction with atorvastatin, 134 later reports have refuted this. 135-137 In a small study of 45 patients randomly assigned to receive either atorvastatin or pravastatin in the background of clopidogrel after ACS, neither statin attenuated clopidogrel-induced platelet aggregation after 5 weeks of treatment. 135

A prospective study of 75 patients undergoing coronary stenting - diabetes in the TRITON-TIMI 38 study, a 30% reduction in major also confirmed the absence of an early interaction between cardiovascular events was observed in patients with diabetes clopidogrel and atorvastatin, according to several measures of treated with prasugrel, in comparison with a 14% reduction in platelet function. ¹³⁶ In fact, patients treated with statins alone had those without diabetes. ¹⁴⁸ There was also no significant difference decreased platelet activity and decreased expression of the thrombin in the rate of major bleeding in patients with diabetes who took receptor protease-activating receptor 1, which support the notion of prasugrel or clopidogrel, which represented a greater net clinical an independent statin antiplatelet effect. ¹³⁶

The drug interaction between clopidogrel and the widely used PPIs has also raised concerns. The mechanism of this interaction is placement with regard to ischemic complications and stent not certain, but it may stem from impaired intestinal absorption of thrombosis. In a subgroup analysis of patients receiving stents in clopidogrel and PPI-induced inhibition of CYP2C19, the major TRITON-TIMI 38, prasugrel conferred 20% and 18% relative enzyme involved in the activation of clopidogrel. 138 An early report reductions in the rate of the primary endpoint among patients highlighted this interaction, demonstrating greater levels of platelet receiving bare-metal and drug-eluting stents, respectively. 149 In reactivity as measured by vasodilator-stimulated phosphoprotein patients with stents, prasugrel was also associated with a 58% (VASP) phosphorylation in patients treated with clopidogrel and relative reduction in stent thrombosis. 149 According to an PPIs. 139 In the Omeprazole Clopidogrel Aspirin Study (OCLA), 124 analysis of STEMI patients in TRITON-TIMI 38, prasugrel was patients receiving coronary stents were randomly assigned in a also more effective than clopidogrel with a 3% absolute risk double-blind manner to receive omeprazole, 20 mg daily, or placebo reduction in the primary endpoint at 30 days. 150,151 in addition to standard clopidogrel and aspirin therapy. ³⁶ With the *Guidelines* use of a VASP assay, a marker of clopidogrel-induced platelet inhibition, the investigators demonstrated greater mean platelet In a focused update of the STEMI guidelines, the ACC/AHA reactivity in patients treated with omeprazole. However, the gave a Class 1 recommendation to the addition of a 300- to 600attenuation of platelet inhibition by PPIs may not be a class effect. In mg loading dose of clopidogrel or 60 mg loading dose of another study of platelet activity in patients treated with clopidogrel prasugrel in all patients with STEMI who were undergoing PCI. and pantoprazole or esomeprazole, neither PPI was associated with 152 In patients receiving fibrinolytic therapy and nonpri mary a change in the mean platelet reactivity index in comparison to PCI, clopidogrel is favored as the thienopyridine of choice, patients taking clopidogrel without PPIs. 140 In a retrospective because of the lack of data for prasugrel in the setting of analysis of more than 8000 veterans treated with clopidogrel after fibrinolytic therapy. 152 Similarly, for patients with a history of ACS, use of PPIs was associated with a greater risk of stroke or TIA, prasugrel is not recommended. Clopidogrel (75 rehospitalization for ACS or death after adjustment for multiple mg daily) or prasugrel (10 mg daily) is recommended for 12 potential confounders (adjusted odds ratio 1.25). 141 Among patients months after placement of a bare-metal or drug-eluting stent. In treated with clopidogrél and PPIs, 14.6% had a recurrent patients scheduled to undergo elective CABG, discontinuation hospitalization for ACS, compared with 6.9% treated with of clopidogrel and prasugrel for a minimum of 5 and 7 days, clopidogrel alone. 141 A more recent observational study from a respectively, is recommended. 152 The ACCP recommended the randomized clinical trial did not demonstrate any clinical addition of clopidogrel, 75 mg daily, to aspirin for patients with interaction, despite ex vivo evidence of a blunting of the antiplatelet symptomatic coronary artery disease. 80 With regard to effect of clopidogrel. 142 The clinical significance of this interaction secondary prevention of stroke, the AHA/ASA suggested that and its contribution to adverse cardiac events was not addressed in aspirin, combination aspirin and dipyridamole, and clopidogrel the context of a randomized controlled trial until as recently as 2009. alone are all reasonable anti platelet strategies. ¹⁵³ Combination In the results of the Clopidogrel and the Optimiza tion of aspirin and dipyridamole is preferred over aspirin alone, Gastrointestinal Events (COGENT) trial, there was no evidence of however. 153 cardiovascular harm from the combination of clopidogrel with proton pump inhibitors. 143

subjects, patients with prior stroke or TIA, and those weighing less NOVEL AGENTS than 60 kg. In a prespecified analysis of patients with and without

Prasugrel

Prasugrel is a newer member of the thienopyridine family with several theoretical advantages over its predecessors ticlopidine and clopidogrel. Although it is also a prodrug, its onset of action occurs in less than 30 minutes, and it has been demonstrated to be 10 times more potent than clopidogrel in animal models. 144 Furthermore, common genetic variants of CYP450 polymorphisms do not appear to be associated with a reduction in the antiplatelet effect of prasugrel. 145,146 In the TRITON-TIMI 38 study, more than 13,000 patients at moderate to high risk with ACS who were undergoing PCI were randomly assigned to receive either prasugrel (a 60-mg loading dose, followed by 10 mg daily) or clopidogrel (a 300-mg loading dose, followed by 75 mg daily) for up to 15 months. 147 Prasugrel was associated with a 19% relative rate reduction in the composite endpoint of cardiovascular death, nonfatal myocardial infarction, or stroke. This finding was counterbalanced by a 32% increase in the rate of major bleeding in subjects who took prasugrel. ¹⁴⁷ A post hoc analysis by the investigators concluded that there was either no net clinical benefit or net harm in three particular subgroups: elderly Although a full discussion of glycoprotein IIb/IIIA (GP IIb/ IIIA) inhibition is not within the scope of this chapter, the historical experience with oral GP IIb/IIIA inhibition is noteworthy. A meta-analysis of the four large, randomized trials of oral GP IIb/IIIa inhibitors that included more than 33,000 patients conclusively demonstrated the deleterious effects of this class of

benefit than in patients without diabetes. 148

Prasugrel has also been shown to confer benefit after stent 7

antiplatelet agents. 154 According to aggregate data, oral GP IIb/IIIa inhibitors were associated with a 31% increase in mortality. 154 Hence, these agents are no longer used for antiplatelet therapy.

Several ongoing trials of novel platelet inhibitors may add to the current therapies for cardiovascular disease. The orally reversible P2Y ₁₂ receptor antagonist, AZD 6140 (ticagrelor), was studied in the Platelet Inhibition and Patient Outcomes (PLATO) trial. 155 Unlike the thienopyridines, it is not a prodrug, and thus hepatic metabolism is not needed to produce an active metabolite. 156 Additional theoretical benefits are rapid onset and offset of action, as well as greater platelet inhibition than with clopidogrel. 157 In a randomized, double-blind study of 18,624 patients with ACS, the PLATO trial demonstrated a decreased risk of the composite endpoint of vascular death, stroke, or myocardial infarction in patients

80 who received ticagrelor, 90 mg twice daily, in comparison with clopidogrel, 75 mg daily (9.8% versus 11.8% respectively). 155 This benefit was achieved without an increase in major bleeding. The results of a phase II study evaluating the efficacy of ticagrelor in patients identified as clopidogrel non- responders are also awaited. ¹⁵⁸ Another reversible P2Y ₁₂ receptor antagonist available in both intravenous and oral forms, PRT060128, is also being compared with clopidogrel in phase II trials of patients undergoing elective PCI (INtra veNous and Oral administration of elinogrel, a selective and 7 reversible P2Y 12 -receptor inhibitor, versus clopidogrel to evaluate Tolerability and Efficacy in nonurgent Percutaneous Coronary Interventions patients [INNOVATE-PCI]) 159 and STEMI (Early Rapid ReversAl of platelet thromboSis with intravenous Elinogrel before PCI to optimize reperfusion in acute Myocardial Infarction [ERASE MI]). 160

Yet another target of platelet inhibition is thrombinmediated platelet aggregation. Thrombin is a potent agonist of platelet aggregation through its interaction with the proteaseactivating receptor 1. 161 The potential incremental clinical benefit of the thrombin receptor antagonist SCH 53038 is being evaluated in two large phase III trials. In the Thrombin Receptor Antagonist for Clinical Event Reduction in Acute Coronary Syndrome (TRA*CER) study, approximately 12,500 patients with ACS will be randomly assigned to receive SCH 53038 or placebo for 1 year, in addition to standard medical therapy. 160 The Thrombin Receptor Antagonist in Secondary of Atherothrombotic İschemic Thrombolysis In Myocardial Infarction 50 (TRA 2°P-TIMI 50) trial is a large, randomized, double-blind, placebo-controlled trial designed to evaluate the efficacy of a 2.5-mg daily dose of SCH 53038 in comparison with placebo in patients with a history of myocardial infarction, stroke, or peripheral artery disease who were being treated with aspirin, clopidogrel, or both. 162 Another novel platelet-activating receptor antagonist, E5555, is being investigated in phase II trials in the Lessons biological activity beyond protease-activating receptor 1 blockade. 164

Cilostazol

diabetes.

With its pleiotropic effects, cilostazol may be added to the follow-up period of 3.5 years, armamentarium of antiplatelet therapy after PCI. In early studies of cilostazol, researchers reported a reduction in intimal proliferation and restenosis after directional coronary atherectomy and balloon angioplasty. 169,170 In a pooled analysis of 23 trials that included more than 5000 patients, cilostazol was found to be associated with a reduction in the risk of both restenosis and the need for repeat revascularization after PCI. 171 More recently, a prospective, randomized trial of triple-antiplatelet therapy - in which diabetic patients who had received drug-eluting stents were given aspirin, clopidogrel, cilostazol – demonstrated reduced angiographic restenosis, as well as target lesion revascularization, in comparison with standard dual-antiplatelet therapy. 172 The protective effect of cilostazol may be partly attributable to attenuation of endothelial senescence induced by drug eluting stents. 173 As might be expected, triple-antiplatelet therapy

results in more potent inhibition of ADP-induced platelet aggregation than does conventional dual-antiplatelet therapy. ^{174,175} Despite these data, the role of triple-antiplatelet therapy in current clinical practice remains uncertain. A retrospective study from a Korean registry provided important insight in this area. In this study of 4203 patients with STEMI who underwent PCI, triple-antiplatelet therapy was associated with fewer major cardiac events, cardiac death, and total mortality than was dual-antiplatelet therapy. 176 The increased mental benefit of cilostazol may prove to be particularly useful in patients with clopidogrel resistance. This hypothesis was tested in the Adjunctive Cilostazol Versus High Maintenance Dose Clopidogrel in Patients with Clopidogrel Resistance (ACCEL-RESISTANCE) study. 177 In this small study of 60 patients undergoing PCI, patients with high post treatment platelet reactivity more than 12 hours after a 300-mg dose of clopidogrel were randomized to receive either 100 mg of cilostazol twice daily or 150 mg of clopidogrel daily. Adjunctive cilostazol was associated with greater platelet inhibition after 30 days than were high maintenance doses of clopidogrel. 177

Dipyridamole

Dipyridamole has a variety of vascular effects that may contribute to from Antagonizing the Cellular Effects of Thrombin Dipyridamole has a variety of vascular effects that may contribute to (LANCELOT) study. 163 In vitro, E5555 also appears to possess its efficacy in cerebrovascular disease, for which it has been widely studied. Dipyridamole inhibits adenosine uptake by red blood cells, which in turn stimulates adenylyl cyclase and subsequent platelet formation of cAMP, an inhibitor of platelet aggregation. 178 An additional antithrombotic effect arises from inhibition of endothelium phosphodiester ase 5, and stimulation of nitric Although first approved in the United States in 1999 for the oxide/cyclic guanosine mono phosphate (cGMP) signaling. 179 treatment of intermittent claudication, cilostazol has been used Dipyridamole has also been shown to exhibit antioxidant and direct as an antiplatelet agent in Asia since 1988. 165 Cilostazol exerts anti-inflammatory effects. 178 Although results of an early its principal antiplatelet effect through selective inhibition of randomized trial of high-dose aspirin and dipyridamole versus phosphodiesterase 3 in platelets and vascular smooth muscle aspirin or placebo were not suggestive of an additional benefit with cells. 165 This leads to increased levels of platelet let cyclic regard to recurrent stroke, 180 subsequent large-scale studies have adenosine monophosphate (cAMP) and ultimately results in provided an evidence base to support its use. The first European inhibition of platelet aggregation and arteriolar vasodilation. Stroke Prevention Study (ESPS) revealed a 33.5% reduction in stroke 166 Antimitogenic effects and inhibition of cAMP uptake may or all-cause death among 2500 patients with a recent stroke or TIA also play a role in the mechanism of action of cilostazol. 166 The who were treated with 75 mg of dipyridamole and 330 mg of aspirin efficacy of cilostazol in the symptomatic management of three times daily, in comparison with placebo. 181 In a 2 x 2 factorial patients with intermittent claudication has been demonstrated design, the European Stroke Prevention Study 2 (ESPS 2) randomly in several trials. In one randomized, placebo-controlled trial assigned 6602 patients with prior stroke or TIA to receive aspirin that included more than 600 patients with intermittent alone (25 mg twice daily), fixed-dose aspirin (25 mg) plus claudication, cilostazol, either 50 mg or 100 mg twice daily, dipyridamole (200 mg) twice daily, dipyridamole alone (200 mg significantly improved pain-free walking distance in twice daily), or placebo. 182 In comparison with placebo, combination comparison with placebo. 167 In a meta-analysis of eight therapy produced a 37% relative risk reduction in stroke, aspirin randomized, placebo-controlled trials that included 2702 alone produced an 18% reduction, and dipyridamole alone patients with moderate to severe claudication, cilostazol produced a 16% reduction. 182 In an open-label design in the resulted in a 67% increase in pain-free walking distance and a European/Australasian Stroke Prevention in Reversible Ischemia 50% increase in maximal walking distance. 168 Furthermore, Trial (ESPRIT), 2764 patients with recent minor stroke or TIA were this benefit was maintained in stratification by gender, age, and randomly assigned to receive aspirin (30 to 325 mg daily) alone or aspirin plus dipyridamole (200 mg twice daily). 183 Over a mean

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^{*}I consider 5 mg if patient weighs <60 kg.

ACS, acute coronary syndrome; AF, atrial fibrillation; ASA-ERDP; aspirin-extended release dipyridamole; CV, cardiovascular; DAT, dual-antiplatelet therapy; MI, myocardial infarction; NSTEMI, non-ST-elevation myocardial infarction; RR, relative risk; RRR, relative risk reduction; STEMI, ST-elevation myocardial infarction; TIA, transient ischemic attack; TLR, target lesion revascularization.

the rate of the primary composite endpoint of cardiovascular death, nonfatal myocardial infarction, nonfatal stroke, or major bleeding was 13% among subjects who took aspirin plus dipyridamole and 16% among patients who took only aspirin. After 5 years of follow-up, 34% of patients discontinued the combination of aspirin and dipyridamole, many because of headache. ¹⁸³The awaited Japanese Aggrenox Stroke Prevention vs. Aspirin Program (JASAP) is

comparing fixed-dose dipyridamole plus aspirin with aspirin, $81\ mg$ daily, for the secondary prevention of stroke. 184

CONCLUSION

Platelets play a fundamental role in thrombosis and inflammation, processes germane to the development of

cardiovascular disease. Inhibition of thromboxane synthesis through aspirin has formed the basis of modern cardiovascular disease prevention. Similarly, ADP inhibition by thienopyridines has proven an essential adjunct in the treatment of patients with ACS, cerebrovascular disease, and peripheral artery disease (Table 7-3). However, despite these therapies, a significant number of patients experience vascular events because of the multiple pathways available for platelet activation . New strategies for platelet inhibition must be developed to achieve greater successes in the treatment of cardiovascular - disease.

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CHAPTER 8

Molecular Biology and Genetics of Atherosclerosis

Paul N. Hopkins

KEY POINTS

- Atherogenesis and contributing factors may be conceptualized in steps, including initiation of endothelial activation and inflammation, promotion of foam cell formation, progression of complex plaques, and precipitation of acute events.
- Redundant signaling pathways lead to endothelial activation in areas of slow flow (especially with flow reversal) and a quiescent endothelial phenotype in areas of higher, unidirectional flow, providing an explanation for predictable locations of atherosclerosis- prone sites.
- Dyslipidemia can lead to endothelial activation through several redundant pathways including LOX-1, TLR4, RAGE, and possibly inadequate protection from HDL.
- Numerous cytokines and chemokines binding to their cognate receptors (many with redundant or overlapping roles) direct accumulation of subendothelial macrophages and other leukocytes.
- Multiple lipoprotein modifications, including nonoxidative, are likely to contribute to foam cell formation in atherosclerotic lesions.
- Knockout studies in mice usually have modest effects on atherosclerosis, perhaps because of redundancy in the pathways.
- The difficulty of demonstrating human genetic associations and their modest effects documented to date (other than the few genes that strongly affect major risk factors) may be explained, in part, by the redundant nature of pathways involved in atherogenesis.

During the past 10 to 15 years, there has been an explosion of knowledge regarding the molecular basis of many diseases, including atherosclerosis. Indeed, the various aspects of atherogenesis illustrate most of the major themes of contemporary molecular biology and cell signaling. Whereas some reluctance to delve into the complexities of these pathways is natural, it is reassuring that many of these signaling pathways and their oddly named members are becoming canonical. Indeed, knowledge of at least some of this newer information will become ever more necessary. This chapter emphasizes the molecular biology and genetics of atherosclerosis, with little discussion of the arguably equally important genetic determinants of the major risk factors.

Perhaps one of the major insights to be gleaned from consideration of the various pathways involved in atherogenesis is their sheer number and complexity. Advances in the science of intracellular signaling 1 and a growing appreciation for the number of cvtokines, chemokines, and receptors involved in cellular communication during atherogenesis led this reviewer to recognize one remarkable feature of such systemsredundancy. Redundancy is seen as a means to ensure the operation of critically important pathways even when one or more elements may be dysfunctional. Thus, a similar element may take over the function of another element or provide a slightly different but overlapping utility. Redundancy is particularly evident in pathways important for survival, on both a cellular and an organismal level. In this regard, atherosclerosis shares numerous pathways involved in defenses against pathogens, inflammation, and cell survival. As it turns out, these pathways are characterized by redundancy (see Table 8-1).

The question then arises: What will be the expected impact of redundancy on our efforts to find genes related to atherosclerosis risk? Whereas the effects on atherosclerosis of many genes may be clearly evident in highly controlled experiments during the short term, their apparent effects may become lost over

time if other mechanisms can substitute for or duplicate their action. If this is the case, redundancy of complex systems may help explain the remarkable difficulty in identifying genes for such complex, common diseases as atherosclerosis and hypertension. Redundancy of the various activation and transduction pathways will be a recurring theme throughout this chapter.

A GENERAL OVERVIEW OF ATHEROGENESIS

Atherogenesis and its clinical expression may be divided into four major steps. Terminology for this model borrows from the cancer literature and was fairly complete in general outline fully 30 years ago. ² The steps include initiation of endothelial activation and inflammation; promotion of intimal lipoprotein deposition, retention, modification, and foam cell formation; progression of complex plaques by plaque growth, enlargement of the necrotic core, fibrosis, thrombosis, and remodeling; and precipitation of acute events, primarily through plaque destabilization and acute thrombosis. Note that acute events may be precipitated by factors unrelated to atherogenesis, for example, through mismatch of arterial oxygen supply and myocardial demand (as with heavy exercise in the setting of uncompensated subtotal coronary occlusion and vulnerable myocardium, leading to ischemia and ventricular fibrillation).

In this scheme, factors frequently act at more than one step of atherogenesis, particularly the major risk factors. For example, elevated lipids can contribute to endothelial activation (with or without oxidation), 3-5 impair nitric oxide (NO) synthesis by endo thelium or its availability, 6,7 lead to foam cell formation (after a variety of possible modifications), 8 increase platelet activation and thrombotic potential, 9 and promote reversible plaque destabilization. 10,11 Obviously , factors that increase thrombotic potential may also help precipitate an acute event if an occluding thrombus ensues.

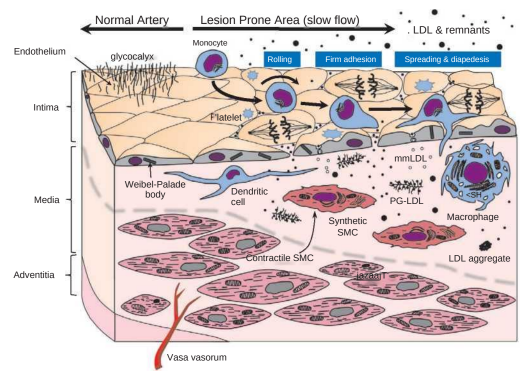


FIGURE 8-1 Initiation of atherosclerosis.

Factors that initiate inflammation are also likely to lead to an unstable plaque. 12

During initiation, several key events occur as illustrated in Figure 8-1. In areas predisposed by hemodynamic factors, atherogenic lipoproteins, including low-density lipoproteins (LDL) and smaller very low-density lipoproteins (VLDL) as well as other remnants of triglyceride-rich lipoproteins (TGRL), infiltrate the intima and are modified by several potential mechanisms. The modified lipoproteins appear to release inflammatory signals to endothelial cells, causing them to become activated. Activated endothelial cells elaborate various chemoattractant and adhesion molecules, such as monocyte chemoattractant protein 1 (MCP-1) and vascular cell adhesion molecule 1 (VCAM-1) recognized by cognate receptors on passing monocytes (and, to a lesser extent, other white cells and platelets) that lead to a sequence of rolling, firm adhesion with spreading and diapedesis or transmigra tion into the subendothelial layer of the intima. Slow or disrupted flow is a major determinant of endothelium activation.

Once in the intima, monocytes transform into activated macrophages capable of ingesting the modified lipoproteins and further amplifying the inflammatory response. Concomitantly, inflammatory cytokines, such as platelet-derived growth factor (PDGF) from adherent platelets and other cells, summon smooth muscle cells to move into the intima, where they change from a contractile to a synthetic phenotype capable of ingesting modified lipoproteins, synthesizing and secreting collagen, and producing various cytokines. Note that platelets can adhere directly to activated endothelial cells, much as white cells do 13; desquamation of the endothelium is not a prerequisite. T cells and mast cells also play an important role in immune signaling and amplification. Thus, during initiation, the intima becomes populated with inflammatory cells poised to do battle with the modified lipoproteins, which are seen as foreign invaders in unauthorized territory.

During the promotion phase (Fig. 8-2), insudation of lipoproteins and their modification continue in proportion to their plasma levels, endothelial permeability, and transarterial pressure gradient that drives fluid convection. Unchecked uptake of remnants or modified lipoproteins by several scavenger receptors on macrophages and smooth muscle cells

leads to formation of foam cells, the hallmark of the growing atherosclerotic lesion. Macrophage foam cells remain capable of relatively rapid egress from the lesion if conditions are favorable (such as a marked reduction in serum lipoprotein concentration) but seem to be retained when lipid levels are high. ¹⁴ Thus, if the balance between lipoprotein entry, foam cell formation, reverse cholesterol transport, and foam cell egress favors cholesterol accumulation, the lesion grows. If excess free cholesterol accumulates within foam cells (particularly with cholesterol monohydrate crystal formation), both apoptosis and necrosis can occur. ^{15,16} This marks the beginning of the formation of the necrotic core as illustrated in Figure 8-2.

As the lipid-rich plaque progresses (Fig. 8-3), accumulated macrophages secrete a host of cytokines and matrix metalloproteinases (MMPs). All migrating cells secrete MMPs to facilitate their diapedesis through the extracellular matrix. Thus, highly cellular plaques would be expected to be more friable. MMPs also undermine the stability of the overlying fibrous cap and contribute to plaque rupture with exposure of the blood to the thrombogenic underlying matrix. In addition, interferon -y (IFN- y), secreted by activated T cells, acts to strongly inhibit collagen formation by smooth muscle cells, further weakening the plaque and fibrous cap. 12 The result can be catastrophic thrombosis and downstream tissue infarction; but more often, there is limited mural thrombosis with subsequent organization leading to saltatory growth of lesions. Other precipitating changes in the plaque include erosions and, importantly, eruption through the endothelium of underlying cholesterol crystals. ¹⁷ Before such episodes of thrombosis, there may be little if any encroachment of the plaque into the arterial lumen because of outward remodeling of the arterial wall to accommodate the growing plaque. Hemodynamic factors and other risk factors seem to influence remodeling and the percentage of the plaque filled with

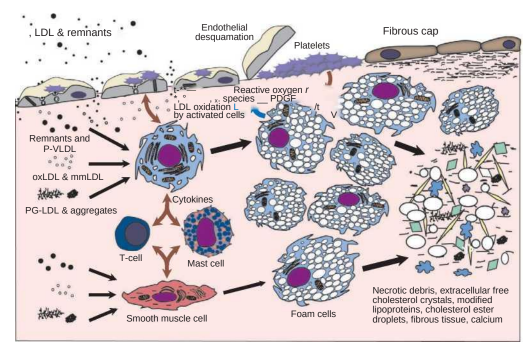


FIGURE 8-2 Promotion of atherosclerosis and foam cell formation.

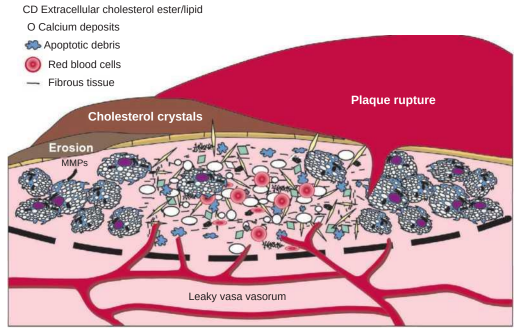


FIGURE 8-3 Progression of atherosclerotic plaques and precipitation of acute events.

fibrous tissue versus lipid. ¹⁸ Calcification appears to be associated with healing or more fibrous plaques, although the overall coronary calcium score remains a powerful predictor of overall atherosclerosis and subsequent risk for coronary events.

When the intimal thickness increases beyond just 0.5 mm, hypoxia induces ingrowth of vasa vasorum. ¹⁹ In recent years, there has been growing appreciation for the potentially major destabilizing effects of these leaky, friable vessels. Vasa vasorum invading lipid-rich plaques remain highly permeable because of constant exposure to inflammatory factors, possibly in a large measure from mast cells. Rather than sudden hemorrhage, a constant leak of red blood cells leads to progressively larger

amounts of red cell markers found in unstable plaques. Thus, red cell membranes may greatly exacerbate the accumulation of free cholesterol and promote the formation of toxic cholesterol crystals. Furthermore, the red cells are a source of strongly prooxidant heme iron, which would favor inflammation and lesion progression in general. Such a scenario may favor progression of lesions for some time even after plasma lipoprotein levels are reduced.



INITIATION OF ATHEROSCLEROSIS

Disturbed Flow and the Atherosclerosis-Prone Endothelial Phenotype

In recent years, endothelial response to flow has been the subject of intense investigation. Sites of atherosclerosis pre dilection occur in areas of low shear or eddy currents (Fig. 8-4) characterized by slow, oscillating (back-and-forth) flow. Such a pattern is referred to as disturbed flow even though the pattern is stable over time in specific areas of the arterial tree. This observation has been confirmed in many observational and experimental studies and is supported by extensive mathematical modeling. ²⁰⁻²² Progression of plaque is predictable in individual human subjects at such locations in the arterial tree. ²³⁻²⁵ Turbulence is virtually never a feature of flow in these sites. ²⁶ Experimentally, atherosclerotic lesions accumulate exclusively in areas of low shear just beyond stenoses created by carotid casts in hyperlipidemic mice. ¹⁸ The unique hemodynamics of the coronary circulation, with near cessation of flow or flow reversal during systole, together with the high pressures generated at the aortic root may explain the marked predilection of coronary arteries to atherosclerosis. Interestingly, rabbit coronary artery endothelial cells expressed only one fifth of the endothelial nitric oxide synthase (eNOS) and greater endothelin-1 (ET-1) compared with aortic endothelial cells. 27 As heart rate increases, relatively more time is spent in systole, when flow is essentially nil. This may help explain the fourfold increase in coronary artery disease (CAD) risk as resting heart rate increased from below 60 to above 100 beats per minute. 28

Multiple, redundant transduction mechanisms endow endothelial cells with exquisite sensitivity to abrupt changes in shear stress. These mechanisms include ion channels, G protein-coupled receptors (GPCRs), the lipid bilayer itself, and the endothelial glycocalyx, with stress transmitted throughout the cell by microfibers and sensed by integrins and at cell junctions. ^{20,29} In general, more rapid flow induces remodeling, which results in larger vessels. Conversely, slow flow results in a decrease in the size of the vessel. In this way, flow becomes a fundamental stimulus for blood vessel development during embryogenesis. ²¹ In addition, flow strongly affects numerous endothelial responses, including endothelial cell migration, mitosis, apoptosis, endothelial layer permeability, inflammation (including white cell adhesion), NO release, and thrombosis. ^{20,21,30}

The pattern of flow is critical in determining the endothelial phenotype. Direct measurement of flow patterns by

SLOW, mildly oscillatory flow with LOW shear stress results in:

· increased cell adhesion molecules

8-5. ³¹ Endothelial cells exposed to the atherosclerosis-prone flow pattern, characterized by slow flow that reversed direction slightly during the course of a cardiac cycle (referred to as oscillatory or disturbed flow), developed a proatherosclerotic phenotype, whereas cells exposed to higher flow rates (higher shear stress or laminar flow) that remained unidirectional although pulsatile had an antiatherosclerotic phenotype. The time course of phenotypic changes and some of the molecular signaling events in the endothelial cells exposed to the respective flow patterns are also shown in Figure 8-5. Platelets and white cells are also much more likely to attach to areas of slow flow, either to endothelium or even when measured for attachment to artificial surfaces such as one coated with E-selectin. ²⁹

Of great importance is the relatively recent recognition that proinflammatory endothelial responses occur only transiently with the onset of flow, after a sudden stepped increase in flow, or with directional change in flow. 29 However, after several hours of continued unidirectional, laminar high shear stress or pulsatile flow (suggesting undulation in the rate of flow but always unidirectional and faster than disturbed, oscillatory flow), these proinflammatory responses are suppressed to below initial (no flow) conditions (see Fig. 8-5). After prolonged exposure (10 to 12 hours) to unidirectional flow, the cells thus display an anti-inflammatory, antioxidant, antiproliferative phenotype, remaining in a relatively quies cent state but active in the production of protective NO and prostacyclin (PGI 2). In contrast, when endothelial cells are exposed to slow oscillatory flow, where direction of flow actually reverses however slightly during the cycle, they never suppress their proinflammatory responses, do not align with the flow, have disorganized cytoskeletons, and develop other features characteristic of lesionprone areas including increased apoptosis, frequent mitoses, greater permeability (particularly at sites of mitosis), shorter glycocalyx, elaboration of cell adhesion molecules, secretion of MCP-1 and endothelin, decreased bioavailability of NO, and increased production of superoxide by NADPH oxidase and heme oxidase. In addition, these endothelial cells increase the production of subendothelial matrix components, such as fibronec tin, which results in enhanced inflammatory responses and a thickening of the basement membrane that may be seen as an increase of intima-medial thickness on ultrasound. ^{20,21,31} The classic inflammatory markers nuclear factor K B (NF- K B) 32 and protein kinase CB (PKCB) 33 as well as other markers of



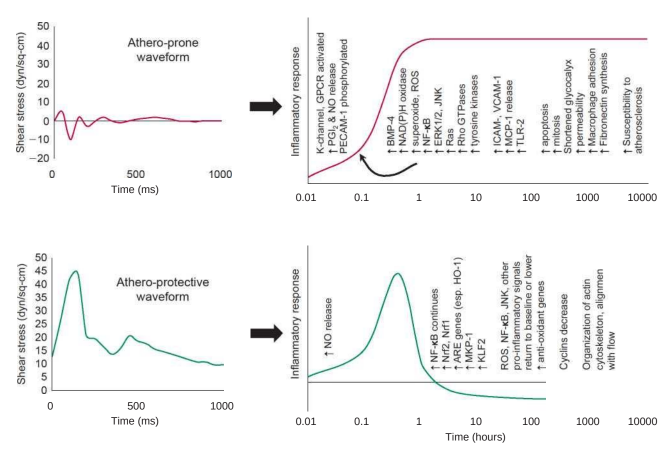


FIGURE 8-5 Hemodynamic characteristics of atherosclerosis-initiating and protective flow and resulting effects

inflammation have been used to track the increased -inflammatory signal seen with early exposure to rapid flow or prolonged exposure to slow flow as well as its suppression with longer term exposure to high shear.

Redundant Mechanotransduction and Cell Signaling Mechanisms

A number of generally proinflammatory signaling pathways are activated immediately or soon after the abrupt onset of flow (Fig. 8-6). If disrupted flow continues (either low shear stress or backand-forth flow), the proinflammatory, proatherogenic state described earlier becomes more fully established. A number of genes in these pathways have been subject to knockout or overexpression in atherosclerosis-prone mice. Although a thorough review of each of the affected signaling pathways cannot be undertaken here, an attempt is made to orient the reader to the location and general function of the specific signaling pathway in the endothelial cell to appreciate the general stage of atherogenesis in which a particular gene may function. Note that whereas a number of the following pathways may be considered canonical and dealt with in general reviews of signaling or in textbooks, they may subserve rather different purposes in specific tissues. The emphasis in the following paragraphs is on such signaling in endothelial cells and in the context of disturbed flow.

The endothelial glycocalyx is important in mechanotransduction, particularly in mediating responses seen in the first few minutes after flow change. The glycocalyx is also involved in inflammatory cell adhesion, thrombosis, migration, and endothelial permeability. The endothelial glycocalyx can be thicker (up to 500 nm or more) than the endothelial cell itself (300 to 400 nm). The glycocalyx is composed primarily of the

proteoglycan syndecan 1 (which consists of a core protein and three or more glycosaminoglycan chains attached). Syn-decan 1 is arranged in a highly structured hexagonal lattice spread over the luminal surface of the endothelial cell with interconnections to the actin cortical web, part of the cytoskeleton. ^{34,35} The stiffness of the proteoglycans and their ability to transmit torque to the endothelial cytoskeleton make for an "exquisitely designed transducer of fluid shear ing stresses." ³⁴ Treatment of endothelial cells with heparinase selectively removes heparan sulfate but not the syndecan 1 core protein ³⁶ and markedly disrupts the glycocalyx layer, reducing many but not all flow-induced responses in endothelial cells, such as NO release in response to shear stress. ³⁷

In the following paragraphs, several pathways of endothelial mechanotransduction are reviewed roughly in the order in which they are activated by the onset of shear stress. They include (1) PKC activation through calcium channels and a GPCR; (2) bone morphogenic protein 4 (BMP4) activation with subsequent generation of reactive oxygen species (ROS) and activation of NF -kB; (3) platelet and endothelial cell adhesion molecule 1 (PECAM-1) and integrin activation; (4) activation of MAPK pathways, particularly JNK; and (5) activation of heat shock proteins.

Within a few seconds after the onset of blood flow, there is an influx of calcium (eg, through the polycystin-2 channel), potassium, and chloride through respective ion channels and activation of GPCRs (such as the bradykinin B2 receptor, which is activated without binding of agonists). The resulting G protein signaling activates phospholipase C with production of diacylglycerol and subsequent activation of PKC § . In addition, there is a sudden initial burst of NO production that is

FIGURE 8-6 Mechanotransduction of shear stress. Onset of flow activates the pathways shown, ultimately resulting in transcription of mostly proinflammatory genes. Exposure to slow or oscillating (back-and-forth) flow results in prolonged stimulation of the pathways shown.

calcium, calmodulin, PKC, and G protein dependent. ³⁸ There may also be a mechanical effect from glypican and CD44 to assist in caveolin-1 phosphorylation and removal of its suppressive effect on eNOS. ³⁹ This stimulation of NO production is generally considered anti-inflammatory and antiatherosclerotic. However, PKC signaling can activate proinflammatory signaling through several MAPK pathways, and knockout of PKC Ş resulted in marked reduction in atherosclerosis. ⁴⁰

BMP4 activation with subsequent activation of NADPH oxidase and heme oxygenase 1 activity together with accumulation of superoxide and other ROS can lead to NF -K B activation. 41 NADPH oxidase may also be upregulated directly in response to the onset of shear stress.

Abrupt onset of flow results in torsional forces transmitted by the glycocalyx to the actin cortical web. This force is thought to be transmitted through actin stress fibers to the dense peripheral actin band and to syndecan 4 and integrins that anchor the endothelial cell to the extracellular matrix. Tractional forces may also be transmitted to lamins on the nuclear envelope. 30 The dense peripheral actin band sur rounds the endothelial cell much like a rubber bumper encir cles a bumper car. 35 It has been proposed that the tractional forces on this band disrupt the weak links between vascular endothelial cadherin (VE-cadherin) proteins of adjoining cells, resulting in the transmission of a signal that is dependent on PECAM-1, VE-cadherin, and vascular endothelial growth factor receptor 2 (VEGFR-2, also called Flk-1). 42 In this process, PECAM-1 appears to act as the true mechanical transducer that directly activates a Src family kinase and also results in the release of bound G a gr. 43 VEcadherin functions as an adapter protein allowing Src to activate VEGFR-2, which then activates phosphatidylinositol 3-kinase (PI3K), which in turn activates integrins as indicated in Figure 8-6. The activated integrins then form new bonds to the extracellular matrix, which triggers activation of Rho, Rac, and Cdc42. In endothelial cells, these work in a coordinated fashion to modify actin polymerization and to mediate cell alignment to flow and other responses. Early changes lead to contraction and increased intercellular permeability. The adapter protein Shc is phosphorylated in response to integrin activation , enabling signaling through other adapters, Grb and Sos, to Ras and Raf with subsequent signaling to the MAPK pathway. Integrin binding and activation also lead to phosphorylation of focal adhesion kinase (FAK), which in turn binds and activates Src, followed by phosphorylation of paxillin and p130 $^{\rm cas}$, which then activates Ras. Alternatively , FAK may first bind Shc with subsequent activation of Src; a third pathway results in activation of Rho. The kinase Src may also directly transmit signals to the MAPK pathway. $^{\rm 21,30,44}$

Yet another result of integrin activation is activation of p21activated kinase (PAK), which may be one of the most important activators of the MAPK pathway resulting from shear stress. This activation appears to depend on PI3K signaling from the PECAM-1, VE-cadherin, VEGFR-2 complex. 45 Not only does PAK activation lead to activated INK (a terminal MAPK), but it may also directly affect various cell junctional adhesion molecules, resulting in increased endothelial permeability. 46 Importantly, the activation of PAK appears to depend on integrin binding to fibronectin in the extracellular matrix. The protein ZO-1, associated with tight junctions, is also involved in the increase in endothelial permeability after the onset of flow or prolonged oscillatory slow flow. Connexin43 (Cx43), unlike other connexins, relocates to cell junctions in response to the onset of shear stress and is increased in atherosclerosis-prone areas of the endothelium. 47

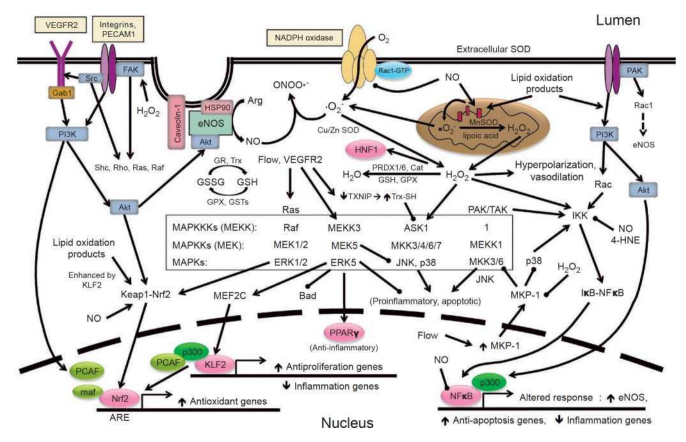


FIGURE 8-7 Pathways leading to the transition from the initial proinflammatory, pro-oxidant state to the quiescent, antiinflammatory, antioxidant state after prolonged exposure to more rapid flow (high shear stress). Lipid oxidation products include 4hydroxy-2-nonenal (4-HNE) and others.

Importantly, there is greater inflammatory response to onset of shear stress when integrins are bound to fibronectin or fibrinogen (characteristically found in extracellular matrix underlying previously injured endothelium) compared with components of the normal extracellular matrix unexposed to prior injury. This difference in inflammatory response seems to be mediated by the binding specificity of different integrins, but activation of Akt also seems to play a role. ⁴⁵The alteration of the extracellular matrix, with increase in fibro nectin, occurs early in atherosclerosis-prone areas as part of the cellular response to disrupted flow, whereas deposition of fibrinogen may occur later. ²¹

As noted before, several of the signals activated on abrupt onset of flow converge on the MAPK/ERK (mitogen-activated protein kinase/extracellular signal-regulated kinase) or simply MAPK pathway, which results, ultimately, in the activation of a variety of transcription factors, including Egr-1 (early growth factor 1) and AP-1 (activator protein 1, which contains c-fos and c-Jun). NF- KB may also be activated by crosstalk with MAP kinases .

The canonical MAPK pathway represents a generic design for a series of three kinases, often held in juxtaposition by a scaffolding protein (with other regulatory proteins sometimes attached, such as IMP or "impedes mitogenic signal propagation"). MAPK designates the terminal kinase. It is activated by a MAPK kinase (MAPKK), also called a MAPK/ERK kinase (MEK). This intermediate kinase is phosphorylated by an upstream MAPKK kinase (MAPKKK or MAP3K) also known as a MEK kinase (MEKK). The MAPK pathway or complex is activated by upstream kinases often in response to tyrosine receptor kinases (classically worked out for EGFR, the endothelial growth factor receptor) or kinases interacting with integrins or other flow sensors. In the canonical EGFR pathway, after binding endothelial growth factor, the

receptor self-phosphorvlates its cytoplasmic domains, allowing binding of one or more adapter proteins (eg, Grb2, which binds directly to the receptor or through another adapter, Shc). Grb2 interacts with the guanine nucleotide exchange factor Sos (son of sevenless), which then activates membrane bound Ras by promoting the exchange of GDP for GTP. Once bound by GTP, Ras is active. In this example (for EGFR), active Ras leads to activation of the protein tyrosine kinase RAF1. RAF1 is actually a MAP3K, the first of a series of three kinases leading to activation of the MAPK ERK. A variety of potential kinases may serve as the initial kinase, MAP3K or MEKK (such as RAF1), as well as multiple potential intermediate kinases and terminal kinases. Mammalian terminal kinases include ERK(1-7), JNK(1-3), and p38(a, Ş, S, and y). Traditionally, ERKs are the terminal kinases for mitogens; p38 and JNK result from stress or cytokine activation. Once activated, terminal kinases can phosphorylate membrane proteins, transcriptional factors in the nucleus, proteins associated with the actin cytoskeleton, various additional signaling molecules, and MAP kinase-activated kinases (MK), which in turn have a series of substrates (such as transcription factors, cell survival, and other signaling proteins). Some examples are given in Figure 8-7. Their control and signal termination are also complex. In endothelial cells, terminal kinases in MAPK/ERK signaling pathways can phosphorylate eNOS and may have various other effects on cell function. Whereas p38 and JNK are generally proinflammatory, ERK5 appears to have multiple anti-inflammatory effects.

Another consequence of early-onset shear stress is the activation of heat shock proteins (HSPs). ⁴⁸ These are generally

considered protective proteins, 49 and recent evidence suggests that an increase in HSP-90, which can act to stimulate eNOS, is brought about not only by prolonged rapid flow but also by statins as a beneficial, pleiotropic effect. 50 HSPs can act as protein chaperones to promote proper folding (especially HSP-70) and other protective functions. HSP-60 may act to chaperone certain cytosolic proteins into the mitochon dria. When cells are not under stress, HSPs are bound to heat shock transcription factor 1 (HSF1) in the cytoplasm. When misfolded proteins are present or the cell is stressed by a number of factors, including ROS, 51 oxidized lipids, or cyto kines, the HSP binds to the misfolded protein (or is otherwise used in the cell), thus becoming separated from HSF1. HSF1 then forms a trimer, moves to the nucleus, and stimulates transcription of HSPs. In the case of HSP-60, levels are clearly upregulated in endothelial cells in atherosclerosis-prone areas of slow flow, with increased expression on the luminal membrane. Cell surface HSP-60 may be important in triggering an autoimmune response that may be important in promoting atherosclerosis (see later). 52

Transition to an Atherosclerosis-Resistant Endothelial Phenotype

After several hours of exposure to laminar flow, the quiescent state described earlier is induced, the actin cytoskeleton - organizes itself into a pattern such that the cells align themselves to be elongated in the direction of the flow with tight cellular junctions, and most cells are found in an arrested state of growth (G $_{0}$ or G $_{1}$). Not only are shear-stressed endothelial cells relatively quiescent, but they become frankly resistant to even potent inflammatory cytokines such as tumor necrosis factor- a (TNF- a). 53,54 The molecular mechanisms that mediate this transition have been the subject of intensive investigation.

Some of the complex intracellular events that appear to underlie the transition from an activated to a quiescent state are depicted in Figure 8-7. Of great importance is the recent recognition that the organized sequence of protective events is at least in part dependent on an initial burst of ROS produced together with increased NO. At the onset of flow, there is a marked increase in production of superoxide anion (-O 2) by NADPH oxidase, possibly mediated by increased activation of the small GTPase Rac (produced by mechanotransduction mechanisms reviewed earlier) or direct effects transmitted through the cell membrane. At the same time, there is a surge in NO production by eNOS due to calmodulin binding and phosphorylation by PKC S. Surprisingly, Cancer appears to be required for normal expression of eNOS as well. 55 Continued and enhanced production of NO is one of the features of the atheroprotective state. This seems to be mediated by a gradual increase in eNOS mRNA as well as by multiple post transcription changes in the eNOS enzyme (such as binding to HSP-90, calcineurin, and certain phosphorylations) that decrease the dependence of eNOS on calcium and calmodulin and increase overall activity. 43

Because -O ₂ reacts rapidly with NO to form peroxynitrite (ONOO-), there is a balance between NO and -O ₂ production that may be significant for signaling events. ⁵⁶ Excess or proprolonged -O ₂ clearly has detrimental effects and can promote cellular damage and apoptosis. ^{57,58} Nevertheless, more controlled release of ROS may be an important signaling mechanism that is key to subsequent protective adaptations by the cell. Early after onset of laminar flow, or with prolonged slow, oscillatory flow, the balance favors -O ₂ production with neutralization of vasodilating effects of NO; after several hours of laminar flow, NO is favored through multiple mechanisms. ⁵⁹ However, some of the vasodilation seen immediately after initiation of flow is mediated by hydrogen peroxide (H ₂ O ₂), which acts as an "endothelial derived hyperpolarizing factor." ⁶⁰ In addition, NO can inhibit mitochondrial electron transport

chain complex I and IV and thereby at least transiently increase mitochondrial production of superoxide anion. 58,61 Lipid oxidation products, such as 4-hydroxy-non-enal, potentially produced during the -O $_2$ spike, are taken up actively into mitochondria, where they also promote -O $_2$ production 58,61 In the cytosol, -O $_2$ is rapidly converted to H $_2$ O $_2$ by copper/zinc superoxide dismutase (Cu/Zn SOD); in the mitochondria, this function is performed by manganese superoxide dismutase (MnSOD). Much of the subsequent "redox signaling" is likely to be mediated by H $_2$ O $_2$, as it is more stable and freely membrane permeable, unlike -O $_2$ or other free radicals. 61,62 Nevertheless, H $_2$ O $_2$, -O $_2$, ONOO- , and hydroxyl radical (-OH) can all contribute to a pro-oxidant environment sensed, in part, through effects on free -SH groups on cysteine residues in various signaling proteins.

Cellular antioxidant defenses depicted in Figure 8-7 include not only the SOD enzymes but catalase, various glutathione peroxidase enzymes, and glutathione- S -transferases (GSTs, which not only convert superoxide to water but reduce a number of other toxic, oxidized molecules by coupling with the GSH to GSSG reaction) as well as glutathione reductase, peroxiredoxins (PRDXs), and thioredoxin (Trx), which regenerate reduced glutathione. Trx also acts as a signaling "gate" by binding (when Trx is in the reduced form) to the MEKK ASK1 (apoptosis signalregulating kinase 1). Thus, ASK1 is a redox-responsive MEKK. ASK1 appears to be the major mediator of TNF-induced JNK activation in endothelial cells. 63 JNK, as noted before, has major proinflammatory and apoptotic effects. When ASK1 is bound by Trx, it is unresponsive to upstream signals, such as those generated from the binding of TNF- a to its receptor. Trx-bound ASK1 is also targeted for ubiquitination and degradation. 64 Trx is highly responsive to ROS and is released from ASK1 upon early exposure to the onset of flow or oxidative stress or during prolonged slow or disturbed flow. In addition, Trx is bound by thioredoxin interacting protein (TXNIP), which inactivates Trx. Importantly, TXNIP is downregulated by prolonged laminar flow through an unknown pathway. 53 Finally, NO can bind to Trx and increase its binding to and inactivation of ASK1. 63

A key, critically protective pathway that is stimulated by flow in endothelial cells and the early ROS burst involves the induction of numerous antioxidant genes. Many of these genes (which include most of the antioxidant enzymes depicted in Figure 8-7) share an antioxidant responsive element (ARE) in their promoters and are activated by the transcription factor Nrf2 (nuclear factor erythroid 2-like related factor 2). Importantly, Nrf2 is activated by both disturbed and laminar flow, but only laminar flow results in increased expression of the antioxidant genes. This was thought to be mediated in part by the oxidized lipid d15-PGJ 2 generated through cyclooxygenase 2 (COX-2). 65 Nrf2 is bound in the cytoplasm by an inhibitor, Keap1 (Kelch-like erythroid-derived cap'n'collar homology-associated protein 1). Keap1 binding targets Nrf2 for proteosomal degradation. Recently, a mechanism was proposed that is dependent on early stimulation of superoxide and hydrogen peroxide by mitochondria, increased early production of NO, and activation of Akt, which all converge to inactivate Keap1 and to release Nrf2 for transport to the nucleus. Effects of oxidized lipids also seem to accelerate the pathway. 61 Direct binding of NO to Keap1 and phosphorylation by Akt appeared particularly important. Critically important for the Nrf2 pathway is the recently discovered effect of KLF2 to prime Nrf2 for greater upregulation of ARE-responsive genes (see later). This interaction, affecting dozens of genes, is one of several major effects promoted by shear stress and KLF2. 66

The early post-flow, pro-oxidant state promotes signaling through ASK1 and promotes activation of inhibitor of K B



94 (IKB) kinase (IKK). IKK phosphorylates IKB, thereby releasing IKB from NF-KB, targeting IKB for proteosomal degradation, and freeing NF -KB TO be transported to the nucleus, where it acts as a transcription factor for NUMEROUS PROINFLAMMATORY GENES . As noted before, NF-KB EXPRESSION IS FREQUENTLY FOUND TO be increased in atherosclerosis-prone regions (see Fig. 8-6). Nevertheless, NF- KB also acts early to support an increase in eNOS synthesis and has antiapoptotic effects. ⁶⁷ This yin-yang behavior is modified in the setting of prolonged laminar flow, in which the proinflammatory actions of NF-KB ARE ALMOST 8 entirely abrogated or "uncoupled" (as activation of genes, especially seen as a marked increase in eNOS drugs. 75 transcription. 68 In addition, there is negative feedback exerted by transport to the nucleus, and inhibits NF- KB binding to DNA. 69

but it also appears to be critical for downregulation of NADPH induced by flow is currently unknown. oxidase subunits after prolonged exposure to laminar flow. was not demonstrated. 70 Interestingly, endothelial expression protective MEK5, ERK5, and, most importantly, KLF2. of the angiotensin type 1 receptor (AT1R) downregulated by shear stress and NO signaling. 71

appears to be another important mechanism whereby the IN ENDOTHELIAL PHENOTYPE protective endothelial phenotype is brought on by laminar flow. As depicted in Figure 8-7, activation of MEK5 by flow Several groups have examined RNA expression profiles of JNK. ^{63,72} Furthermore, MEK5 activates ERK5, which interferes the transcription factor MEF2C (myocyte enhancer factor 2C), thereby inactivates the proapoptotic factor Bad.

Numerous studies have pointed to the induction of KLF2 as a critical step in the conversion to and maintenance of an atheroprotective state. 73-77 KLF2 is expressed almost exclusively in areas of the vasculature protected from atherosclerosis, whereas it is nearly absent in atherosclerosisprone areas and is clearly upregulated by laminar flow. It is also important in embryonic vascular development. The molecular effects of KLF2 are protean and include: inhibition of activating transcription factor 2 (ATF2), which can be one of the heterodimeric components of AP-1, the key product of the MAPK p38 and a required activating factor for many proinflammatory effects of NF- K B 78; induction of inhibitory Smad7, which blocks transmission of proatherosclerotic signals through the transforming growth factor- \$ (TGF- \$) receptor, together with reduction of nuclear c-Jun, a second component of AP-1, thereby further blocking many

inflammatory signals 79; enhanced transcription of eNOS and the enzyme dimethylarginine dimethylaminohydrolase, which degrades the eNOS inhibitor ADMA (asymmetric dimethyl transcription arginine); increased of thrombomodulin (TM); inhibition of ET-1 and MCP-1; and other major effects. In one study, KLF2 decreased the expression of the following genes by 80% to 90% after exposure of cells to IL-1: IL-1a, IL-1, IL-6, IL-8, IL-15, MCP-1, E-selectin, TNF-a, CXCL10, CXCL11, IFN-y, COX-2, and CCL5. 75

An important mechanism to upregulate KLF2 involves activation seen by a marked suppression of VCAM-1, E-selectin, and IL-8 of p300/CBP-associated factor (PCAF) by a PI3K-dependent but production in response to TNF -a), leaving the cytoprotective effects Akt-independent mechanism. 80 Activated PCAF acts together with of NF-KB (such as an induction of MnSOD and eNOS) intact. 54 p300 to acetylate histones and greatly increase transcription of KLF2-Recent insights into the mechanism of this transition suggest regulated genes. Increased PCAF activity is also related to induction activation by Akt (also activated early after the initiation of flow) of COX-2 independent of KLF2. Thus, PCAF activation appears to be a histone acetylator, p300 (generally referred to as p300/CBP; CBP yet another critical mechanism linking early signaling events to the refers to the p300 homologue CREB-binding protein). p300/CBP onset of flow. Interestingly, statin drugs are potent inducers of KLF2, may also be activated by a short burst of oxidant stress. The activated possibly by way of Akt activation, a finding that lends credibility to p300 acts as a coactivator with NF-KB and greatly affects NF-KB the potential importance of so-called pleiotropic effects of these

MAPK phosphatase 1 (MKP-1) is another gene found to be NO at several steps of the NF - KB pathway. Thus, nitrosylation expressed exclusively in arterial regions protected by high laminar inhibits IKK, stabilizes I K B, induces I K B mRNA, decreases NF- K B flow. 81 As MKP-1 deactivates p38 and JNK by phosphory lating these terminal MAP kinases, it also has the potential for major anti-Not only does NO provide feedback inhibition to NF-KB, inflammatory effects. The precise mechanism by which MKP-1 is

In summary, prolonged flow establishes a quiescent endothelial Thus, when endothelial cells were subjected to shear forces in phenotype by the following redundant and somewhat interacting a cone-and-plate viscometer, NO and -O 2 - production were mechanisms: (1) use of the initial pro-oxidant burst as a trigger for upregulated 6-fold and 2.5-fold, respectively, by 2 hours. There the induction of cellular antioxidant defenses; (2) capitalizing on the was no change in NADPH oxidase subunit number at this reduction of ROS to down-regulate JNK (through decreased TXNIP point. It is possible that a higher earlier peak for -O 2 was and increased Trx) and NF-KBSIGNALING; (3) activation of Akt/PKB, missed. Afterwards, -O₂-declined until it was about 50% of the resulting in strong eNOS upregulation and antiapoptotic signals as initial level after 24 hours of laminar shear stress while NO well as diversion of NF-KB TO a largely anti-inflammatory role with production had increased markedly. The investigators found further support of eNOS induction; (4) greatly upregulated NO that there was an approximate 50% decrease in the expression production, resulting in inhibition of NF-KB inflammatory signaling, of the activity-limiting gp91 phox subunit of NADPH oxidase downregulation of NADPH oxidase activity, and decreased while eNOS protein expression was increased 3.5-fold. The expression of the AT1R as well as other protective activities; and (5) reduction in gp91 phox expression was shown to be NO MAPK pathway crosstalk and induction of MKP-1, leading to dependent, although the specific signal transduction pathway inhibition of p38 and JNK signaling while promoting signaling by

Crosstalk between different MAPK pathway kinases GENOMIC STUDIES OF FLOW-MEDIATED CHANGE

inhibits activation of the proapoptotic, inflammatory MAPK endothelial cells exposed to the atherosclerosis-prone versus protective flow patterns. 31,66,73,78,82-86 In these studies, expression of with signaling downstream of JNK. 63 Perhaps even more up to 10,000 or more genes could be assessed simultaneously by important is the induction of KLF2 (Kruppel-like factor 2) by examining differences in the levels of mRNA found in harvested endothelial cells exposed in vitro to the different flow patterns. which is activated by ERK5. ERK5 also phosphorylates and Although there were a number of differences in technical approaches used in these studies and differences in specific findings of which genes were significantly upregulated or downregulated, this genome-wide approach generally confirmed the findings of candidate genes and molecular studies in which genes associated with inflammation and susceptibility to atherosclerosis tended to be increased in cells exposed to the slow-flow, oscillatory pattern, whereas such genes were suppressed in the cells exposed to rapid, unidirectional pulsatile flow.

In one of the studies, endothelial cells were harvested from

of aorta from normal pigs. 85,87 This in vivo approach suggested that whereas proinflammatory genes were frequently increased in the atherosclerosis-prone areas, some potentially compensating genes were also upregulated, including several antioxidant genes, such as glutathione peroxidase. Also, there seemed to be little evidence of frank inflammation in these arteries. Thus, active NF- K B was not found in greater levels in nuclei, nor were several key adhesion receptors found to be expressed at higher levels on the cell surface. The authors interpreted these results to suggest that disrupted or slow flow primes the endothelium to respond with an enhanced inflammatory response if it is exposed to additional risk factors. Thus, "a delicate balance of pro- and antiatherosclerotic mechanisms may exist simultaneously in the endothelium of lesion-prone sites of the aorta to create a setting of net vulnerability to atherogenesis, but with protective measures also present. A shift from atherosusceptibility to atherogen-esis may occur by inhibition of the protection." 87

This concept of predisposition is supported by the finding of an increase in both bound NF-KB AND its inhibitor IKB IN THE cytoplasm of endothelial cells from atherosclerosis-prone areas of the aorta in mice but with no increase of NF- KB IN the nucleus. Nevertheless, there was much greater activation of NF- KB and its inducible gene products in these same hemodynamically prone areas after the mice were treated with lipopolysaccharide or after feeding LDL receptor (LDLR)-deficient mice a high-fat diet to induce hyperlipidemia. 88

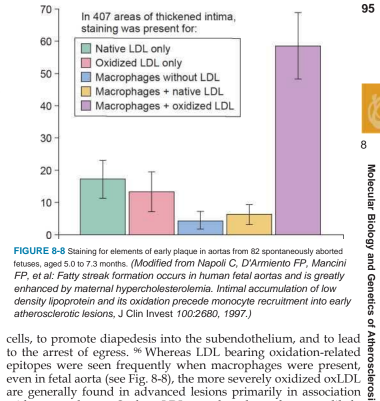
Further Endothelial Activation by Dyslipidemia

The well-known observation of monocyte adherence to endothelium are generally found in advanced lesions primarily in association & in atherosclerosis-prone areas just days after induction of severe with macrophages. Such oxLDL are thought to be more likely hypercholesterolemia by cholesterol feeding in animals can be better formed by exposure to myeloperoxidase-produced hypochlorous appreciated in light of the above molecular mechanisms. 89,90 Factors acid from activated macrophages. 57,100 This consideration raises the that even modestly activate endothelial cells would be expected to question of whether oxLDL help summon the macrophages initially initiate sub stantially greater effects in the already "primed" cells or whether oxLDL are formed only after activated macrophages found in atherosclerosis-prone areas. 88 Hyperlipidemia clearly - have arrived. Nevertheless, even if formation of oxLDL or provides one or more signals for such activation, 91 whereas low - chlorinated and nitrosylated forms of LDL require preexisting ering of serum cholesterol, even if only by dietary means, clearly macrophages, such modified LDL could subsequently perpetuate decreases endothelial activation. 7,92 As early as 2 hours after endothelial activation and otherwise promote atherosclerosis. 101 injection of human LDL into rabbits, grapelike clusters of aggregated One important issue to note in this regard is the near absence of LDL could be seen enmeshed in focal areas of the subendothelial myeloperoxidase in experimental mouse atherosclerotic lesions, matrix. 93 VCAM-1 and MCP-1 are expressed by endothelial cells suggesting a major species difference with humans and providing a within at least 3 weeks of starting a high-cholesterol diet in rabbits. likely explanation for the lack of effect on murine atherosclerosis 92 Not only do lipo proteins preferentially accumulate in after knockout of myeloperoxidase, whereas humans with myelo atherosclerosis-prone areas, but hyperlipidemia itself clearly peroxidase deficiency appear to be protected from atherosclerosis. 98 increases the permeability of the endothelium and total area of Perhaps the strongest evidence in favor of the oxidized LDL susceptibility, suggesting direct activation. 89,90 Both activation of hypothesis comes from the decrease in atherosclerosis seen after endothelium with increased cell turnover and decreased glycocalyx immunization of animals with various forms of oxidized LDL or height appear to mediate this increased permeability due to malondialdehyde-treated LDL. 102-104 Still, the immunization hyperlipidemia. 94 This appeared true with even modest appeared to be most effective at later stages of atherosclerosis. 103 hyperlipidemia as newly synthesized, radiolabeled thymi dine (an Given these considerations, there remains the need to explain how index of cell turnover) in aortic endothelial cells tripled in just 3 days the endothelium is activated to attract the monocytes in the first after starting a high-cholesterol diet in pigs when serum cholesterol place. levels had increased to only 187 mg/dL (normal, 74 mg/dL). 95 Thus, hyperlipidemia clearly activates the endothelium, but the precise physiological exposure to copper or iron ions), endothelial and signals mediating this effect have been less easily identified. 96,97 smooth muscle cells can mildly oxidize LDL in culture, and Given the current evidence, it would appear that multiple mechanisms linking hyperlipidemia to endothelial activation are likely, further illustrating the principle of redundancy. Several of these possible mechanisms are presented briefly here.

LDL Oxidation

Much evidence has been forwarded in support of the LDL oxidation hypothesis for endothelial activation (and still

more for foam cell formation, to be discussed later), 96,98 Native LDL appears to accumulate in the intima of human fetuses before macrophages are found. Appearance of oxidized LDL seems to follow native LDL, and the oxidized LDL are frequently present without macrophages nearby. However, the most common finding was the presence of intimal macrophages together with oxidized LDL (Fig. 8-8). 99 Severely oxidized LDL (oxLDL) have been shown to stimulate adhesion of both macrophages and T cells to endothelial



Aside from oxLDL (usually produced in vitro with non -

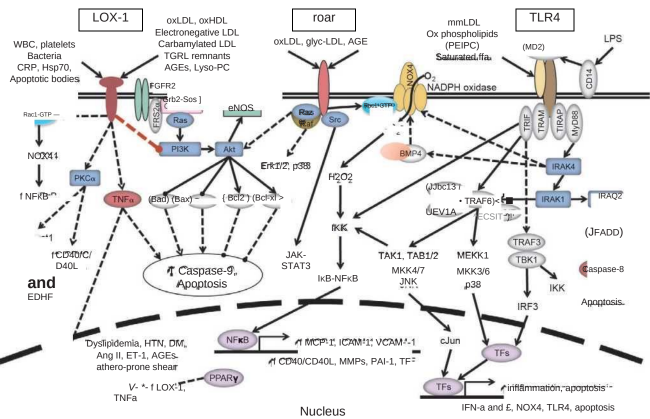


FIGURE 8-9 Activation of endothelial cells through LOX-1, RAGE, or TLR4 signaling. Highly parallel pathways are present in macrophages that can lead to activation and foam cell formation.

such "minimally modified" LDL (mmLDL) can also activate endothelial cells and initiate monocyte adherence and trans migration. These events were prevented by the presence of highdensity lipoproteins (HDL) or antioxidants. 105 In these studies, only monocyte, not neutrophil, adhesion and migration was stimulated, much as might be expected for atherosclerosisrelated endothelial activation. Presumably, LDL trapped in the subendothelial space would be exposed to sufficient ROS from activated endothelial cells to result in such minimally oxidized LDL. Formation of mmLDL or oxLDL in the plasma is thought to be unlikely because of potent antioxidant factors there. Some have questioned whether even mmLDL is formed in vivo in the 57 However, surprisingly antioxidant-rich intima. observations in human fetuses cited before would suggest that at least some oxidation is possible and occurs before the arrival of macrophages.

mmLDL and Toll-Like Receptor 4 (TLR4)

Perhaps the strongest evidence for a pathway mediating endothelial activation by mmLDL is by way of toll-like receptor 4 (TLR4). 106-108 TLR4 is a major mediator of innate immunity and the main response receptor for bacterial lipopolysaccharide. A number of pathogen-associated molecular patterns (PAMPs) appear to trigger inflammatory responses through TLRs. Oxidized or otherwise altered phospholipids protrude abnormally (much like whiskers) from the cell membrane and trigger innate immune receptors and natural antibodies that act to clear the damaged phospholipids. 109,110 These considerations may be most relevant to macrophage activation and foam cell formation, considered later. Nevertheless, endothelial cells from C3H/HeJ mice, which have an inbred mutation in TLR4, are essentially unresponsive to mmLDL and are protected from dietinduced atherosclerosis . 111 Furthermore, mmLDL as well as associated oxidized phospholipids were shown to transmit signals to activate NF-KB through TLR4. 106 Knockout of TLR4 or

its downstream adapter protein MyD88 (myeloid differentiation factor 88) resulted in unresponsiveness to mmLDL in endothelial cells and decreased atherosclerosis. ¹⁰⁸ As noted later, TLR4 may also be involved in other mechanisms of endothelial activation in relation to atherosclerosis. Several features of TLR4 signaling are shown in Figure 8-9. TLR2 may also be involved in endothelial activation. ¹¹²

Role of the Oxidized LDL Receptor 1 (LOX-1)

with oxLDL powerful Early studies demonstrated proinflammatory and apoptotic effects on incubation with endothelial cells. Subsequent studies identified LOX-1 (lectinlike oxidized LDL receptor 1), coded by the OLR1 gene (oxidized LDL receptor 1), as a scavenger-type receptor expressed on the surface of endothelial cells that likely mediated these effects. LOX-1 is also expressed on monocyte-macrophages and platelets, but unlike macrophages, LOX-1 is the only scavengertype receptor expressed highly on endothelial cells. LOX-1 may be considered another PAMP-recognizing receptor with a key role in innate immunity. 109 Importantly, LOX-1 is not expressed constitutively but is induced by most of its ligands, including oxLDL, as well as by factors that upregulate NADPH oxidase, such as angiotensin II. Besides oxLDL, LOX-1 is bound and activated by gram-positive and gram-negative bacteria, apoptotic bodies, senescent red blood cells, activated white blood cells and platelets, advanced glycation end products (AGEs), lysolecithin, and, recently recognized, C-reactive protein. 113,114 LOX-1 is expressed particularly in atherosclerosis-prone areas or over atherosclerotic plaque and has been reported to be increased by

hyperlipidemia, diabetes, and hypertension and after exposure to various cytokines. 115-117

Details of LOX-1-mediated intracellular signaling continue to be worked out. Some of the identified pathways are depicted in Figure 8-9. Intracellular signaling pathways activated by LOX-1 include RhoA (resulting in eNOS downregulation); a Rac1mediated burst in NADPH oxidase activity 118; the Ras, Raf, ERK1/2 cascade, leading to increased PAI-1 expression 119; and increased expression of CD40 and CD40L (CD40 ligand) through PKC a signaling, which may act in an autocrine fashion through CD40L signaling to further upregulate several inflammatory genes including TNF- a and P-selectin. 120 Importantly, activation of LOX-1 leads to inactivation of PI3K. In endothelial cells, PI3K is strongly protective as it normally upregulates eNOS and inhibits apoptosis. Upregulation of PI3K is key to the protective autoregulation by fibroblast growth factor 2 (FGF2) through its own receptor, FGFR2, as shown in Figure 8-9. Therefore, by blocking PI3K and through increased TNF- a production, apoptosis is greatly accelerated. 121,122

If oxLDL were the only lipoprotein to bind to the LOX-1 receptor, its relevance to early endothelial activation would not be so readily apparent for reasons noted before. However, studies note that in addition to oxLDL, LOX-1 is bound and strongly activated by electronegative LDL. ¹²¹ This finding is consistent with a substantial literature demonstrating the endothelial activating and apoptotic potential of electronegative LDL. These particles can be formed by a number of nonoxidative mechanisms as well as by oxidation, including glycation, enrichment with nonesterified fatty acids, treatment with cholesteryl esterases, phospholipase A 2, platelet activating factor acyl hydrolase (PAF-AH, also called LpPLA 2), and, importantly, sphingomyelinase. 123,124 Electronegative LDL are found in a much higher proportion of plasma LDL than are oxidized LDL (up to 10% of normolipidemic LDL). Recently, evidence was presented that LDL themselves carry a sphingomyelinase activity that promotes formation of electronegative LDL and that promotes LDL aggregation as seen subendothelial, trapped LDL. 125 Furthermore, sphingomyelinase and other enzymes capable of modifying LDL and promoting aggregation are present in the artery wall and likely promote aggregation of LDL. 124 Importantly, other studies identify LOX-1 as a receptor that can mediate inflammatory responses to TGRL (triglyceride-rich lipoprotein) remnants. 126

Particles that bind to LOX-1 appear to be relevant in vivo because overexpression of LOX-1 in the liver to remove such particles from plasma led to virtual arrest of atherosclerosis progression. ¹²⁷ Knockout of LOX-1 reduced atherosclerosis approximately 50% in a high-cholesterol/fat-fed LDLR knockout model and preserved endothelial function. ¹²⁸ Over expression of LOX-1 in coronary arteries in apo E knockout mice leads to an atherosclerosis-like vasculopathy. ¹²⁹

TGRL remnants may be particularly atherogenic. In one study, TGRL appeared to activate endothelial cells without any need for modification. 5 In another study, apo C-III, which accumulates on TGRL, appeared to activate PKCB and thereby inhibit protective Akt signaling in endothelial cells. ¹³⁰ In a gene expression study comparing VLDL with oxidized VLDL (oxVLDL), different endothelial pathways were activated. 131 Thus, native VLDL upregulated ERK1/2 and NF-KB MODESTLY WITH little effect on ROS generation, whereas oxVLDL resulted in substantial stimulation of the MAPK p38, NF-KB, and marked increase in ROS production as well as evidence for decreased viability. Later studies by this same group showed much greater endothelial activation by postprandial TGRL after a cream meal compared with fasting TGRL from hypertriglyceridemic patients. In addition, flow-mediated dilation of the brachial artery was impaired by the fatty meal. Post prandial TGRL resulted in upregulation of p38 MAPK, CREB, NF-AT, NF- KB, VCAM-1, PECAM-1, ELAM-1, ICAM-1,

P-selectin, MCP-1, IL-6, TLR4, CD40, ADAMTS1, and PAI- $\bf 97$ $1.^{132}$ The receptors or other mechanisms mediating these responses were not identified.

present on endothelial cells and, like TLR4 and LOX-1, can activate endothelial cells by overlapping intracellular signaling pathways (see Fig. 8-9). JNK activation seems to be particularly important in this regard, but RAGE also activates the JAK-STAT3 pathway. 133 RAGE is generally considered in the context of diabetes-related modifications to proteins or LDL. Indeed, it recognizes a ligand generated by nonenzymatic reactions 8 between glucose and protein lysine residues (carboxy-methyl lysine). Certainly, ligation of RAGE by this ligand, generated in proportion to degree of hyperglycemia, is one means whereby diabetes contributes endothelial to activation atherosclerosis, as demonstrated by use of soluble RAGE competition ¹³⁴ as well as diabetic RAGE knock out models. ¹³⁵ However, hyperlipidemia without diabetes also generates a substantial load of ligands for the RAGE. Indeed, 52% reduction in atherosclerosis was seen in nondia betic, apo E knockout mice that were also deficient in RAGE, whereas those expressing endothelial cell-specific dominant negative RAGE mutations had more than 70% reduction. ¹³⁶ Ligands for RAGE present in oxLDL identified in this model include proinflammatory S100/calgranulins and HMGB1 (high-mobility group box 1). Incubation with S100 activated JNK with elaboration of VCAM-1 in cultured wild-type endothelial cells, whereas activation was substantially reduced in the RAGE knockout or dominantnegative cells. 136 Thus, RAGE represents yet another pathway for endothelial cell activation.

REFLECTIONS ON ENDOTHELIAL ACTIVATION

One of the major messages in these discussions is the redundancy of pathways potentially leading to endothelial cell activation. For modified lipoproteins alone, activation may occur by way of TLR4, LOX-1, or RAGE. The TLR4 and RAGE responses may require oxidative changes, as found in mmLDL or oxidized phospholipids, but may also be triggered by saturated fatty acids (for TLR4); the LOX-1 response appears to include a broader array of triggers. Other potential pathways resulting in endothelial activation from various lipoprotein moieties are also likely. Thus, an intervention that focuses on only one of these pathways, such as oral antioxidants (even acknowledging the limitations of the antioxidants tested thus far), might be expected to be ineffective in reducing coronary events. This can be expected even if oxidation is causally related to disease because of the redundancy of lipoprotein modifications that can lead to endothelial activation. For the same reason, effects from single genes in these pathways may be difficult to detect. Indeed, not shown in Figure 8-9 are the numerous cytokine-mediated pathways that lead to activation. Additional pathways are illustrated by numerous knock out models reviewed in Tables 8-1 and 8-2.

Redundancy seems to be the rule for pathways vital to the survival of the organism. In the case of endothelial activation, the key role such activation plays in defense from microorganisms as well as in clearing the bloodstream of senescent or apoptotic debris seems obvious. In this sense, it may be disingenuous to describe the state of endothelial activation associated with various risk factors as "dysfunction." Indeed, the cells seem to be performing admirably and as expected in response to perceived threats.

TABLE 8—1 Effects on Atherosclerosis of Gene Knockout, Transgenic Expression, or Other Genetic Manipulations Involving Endothelial Cell Activation and Other Early Steps in Initiation *

genes	Effect +	Gene Function
PKC p i/2		A conventional proinflammatory PKC
Egr-1		
		Proinflammatory transcription factor activated by JNK
PECAM-1	— to	Flow sensor, involved in intercellular junction
Fibronectin, EIII	Α	Extracellular matrix component, proinflammatory
Cx37 (BMT)	Т	Regulates macrophage adhesion to endothelium
Cx43		Promotes leukocyte accumulation
eNOS	Т	Synthesis of NO
p47 ^{pl} =		Critical component of NADPH oxidase
Renin		Produces angiotensin I from angiotensinogen
AT1R		Angiotensin receptor 1 (can activate NADPH oxidase)
HO-1	tut	Important cellular enzymatic antioxidant
MnSOD	tut	Important cellular enzymatic antioxidant
PRDX1	T	Important cellular enzymatic antioxidant
LIAS	Т	Synthesis of lipoic acid, a mitochondrial antioxidant
NEMO	-	Component of IKK that phosphorylates I K B
IKB(DN)		Sequesters NF- K B in the cytoplasm until phosphorylated
TLR2	-	A toll-like receptor—normally recognizes PAMPs
TLR4		
		A toll-like receptor, binds modified lipoproteins
MyD88		Adapter protein critical for TLR signaling
LOX-1,		
OLR1 roar		Activation of endothelial cells by modified lipoproteins
Todi		Activation of endothelial cells by modified lipoproteins
PARP-1		Activation of endotherial cells by modified apoproteins
		DNA repair and stress enzyme-activated DNA damage
ROCK1		Transmitted contraction signals from Rho
РІЗКγ , р110γ		Regulates macrophage chemotaxis
5-LO		Rate-limiting enzyme for leukotrienes
		synthesis

^{*}In this and the following tables, studies were whole-body knockout of the gene unless otherwise indicated, performed in either apo E-deficient or LDL receptor knockout mice.

Molecular Biology of Leukocyte Transmigration into Intima

The first step of capturing passing leukocytes has generally been considered to be mediated by endothelial cells expressing on their surface the adhesion molecules P-selectin and E-selectin. Binding of these selectins to their cognate

		ponding Cognate Receptors Cell- Leukocyte Interactions	
Endothelium	Effect	Leukocyte	Effect
Tethering or Rolling			
E-selectin	-	ESL-1, CD44	
P-selectin	-	PSGL-1	-
PSGL-1, CD24, CD34	-	L-selectin	_
Activation by Chemoki M-CSF	nes/Chemoa	ttractants CSF1R	
MCP-1 (CCL2)	— to	CCR2	to
CCL5 (RANTES)	-to	CCR1 CCR5	TT
CXCL1 (GRO a)	-	CXCR2 (BMT)	
CXCL2 (GRO P) CXCL3 (GRO Y) CXCL5-7	_	CXCR2	
CXCL4 (PF 4)		CXCR3B	_
CXCL8 (IL-8)	_	CXCR2	
CXCL9, 10, 11	-	CXCR3	NS
CXCL12 (SDF-1)	_	CXCR4	Т
CXCL16	Т	CXCR6	-
CX3CL1 (fractalkine)		CX3CR1	
CD40	NS	CD40L (CD154)	
Slow Rolling and Firm VCAM-1	Adhesion -	α4β1 integrin (_{VLA4)}	_
Fibronectin		« ₄ P _. , « ₅ P _.	_
ICAM-1	NS to	a _L p ₂ integrin (LFA1)	-
E-selectin	-	» мр ₂integrin	_
VWF	_	O -iib P a integrin	_
Thrombospondin-1 (VN)	_	a vp ₃ integrin	tut
MADCAM1	_	» 4p7 integrin	_
CD137	_	CD137L	_
Transmigration JAM-A	_	LFA1, JAM-A	_
JAM-B		VLA4, JAM-C	
JAM-C (also in smooth muscle cells)	-	MAC1, JAM-B	_
CD99	-	CD99	_
PECAM-1	-to	PECAM-1	-to
ICAM-2	_	MAC1, LFA1	_

[·] In most cases, monocyte interactions are shown. Interactions with other leukocytes and platelets may follow similar steps. Cells other than endothelial cells may produce the chemokines shown.

receptors on leukocytes is relatively loose, with breaking and reforming of the bonds, thereby leading to partial tethering and rolling of leukocytes under the shear forces of the blood circulation. L-selectin on virtually all leukocytes can recognize several markers on activated endothelial cells as well.

[^]Effects on atherosclerosis (for all tables): -, <50% reduction; --, 50%-80% reduction; ---, >80% reduction; T, <100% increased; TT, 100% or greater increase.

BMT, bone marrow transplantation; DN, dominant negative.

BMT, bone marrow transplantation; NS, not significant; —, not tested.

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Tethering and rolling bring the leukocyte into intimate contact with the endothelial surface, where the leukocyte encounters chemotactic cytokines or chemokines such as MCP-1 (also referred to as CCL2) and monocyte colony stimulating factor (M-CSF) as well as other surface-immobilized chemokines that are held close to the endothelial surface by proteoglycans in the glycocalyx. More than 40 chemokines are known, again illustrating the redundancy of key systems. ¹³⁷

The current terminology for the chemokines refers to characteristically conserved cysteine residues in the N terminus. These cysteines can have no intervening amino acids (CC chemokines), one separating amino acid (CXC chemokines), or three intervening amino acids (CXXXC or CX3C chemo kines); L refers to the chemokine ligand and R to the receptor. For example, some of the surface-immobilized chemokines noted before include CXCL1, CXCL2, CXCL4, and CCL5 in addition to MCP-1 (CCL2). Chemokines presented by activated endothelial cells bind to cognate receptors that are all GPCRs (such as CCR2, the receptor for MCP-1) located on the surface of the leukocyte. This binding leads to further activation. At least 20 such receptors are known. Even before this step, rolling appears to promote movement of these GPCRs from intracellular sites to the cell surface in association with cholesterol-rich rafts. One study demonstrated the presence of MCP-1, GRO a , and IL-8 on the endothelium of human athero sclerotic plaque that was capable of inducing attachment and spreading of test monocytes, suggesting the relevance of prior, mostly mouse-based models and further illustrating redundancy in this early step of

Binding of chemokines to their cognate GPCRs initiates "inside-out signaling," with the result being an alteration in the extracellular configuration of leukocyte integrins leading to enhanced binding affinity to endothelial VCAM-1 and ICAM-1 (or other ligands). This enhanced interaction then leads to slow rolling, adhesion strengthening, spreading, and firm adhesion. The firm binding of leukocyte integrins with their ligands then initiates "outside-in signaling" and clustering of the integrins into focal adhesions. Outside-in signaling refers to a host of integrated signals initiated by the bound integrins that result in intraluminal crawling as the leukocyte seeks an opportune site for penetration and finally diapedesis or transmigration of the leukocyte into the subendothelium. This may occur by passage of the leukocyte between endothelial cells by interactions with various gap junction proteins (paracellular migration) or directly through thinned segments of endothelial cells (transcellular migration). For monocytes, this is followed by transformation into tissue macrophages or dendritic cells. There are at least 24 different integrin heterodimers that mediate this signaling on various leukocytes. 139

Each progressive step of this process is associated with greater activation of the leukocyte and coordinated intra cellular and intercellular signaling with endothelial cells. Whereas monocyte transmigration has been considered to predominate in atherosclerotic lesion formation, the participation of other leukocytes and subsets of monocytes and T cells has recently been more fully recognized. 140 Platelets can also adhere to activated endothelial cells and may aid in the binding of other white cells. T cells, mast cells, and even a few B cells and neutrophils also find their way into the initial lesion. Once they are in the subendothelial space, there is an exchange of cytokines between the various activated cells that further amplifies the inflammatory response. The molecular biology of these steps in relation to atherosclerosis has been extensively studied and is the focus of several recent, excellent reviews. 13,140-144 Some of the ligands and receptors mediating these endothelial cell-leukocyte interactions are listed in Table 8-2. However, virtually any presentation of leukocyte activation and interaction with the endothelium must be considered a simplification. One review listed 47 proteins that were thought to be regulatory for at least 900 total proteins and 6000 protein-protein interactions . 145 Gene expression profiling identified 400 genes that were upregulated

or downregulated by at least twofold after exposure of endothelial cells to IL-1 \S , IFN- y , and TNF- a . 146 About 600 to 1000 genes were similarly regulated during the transition of monocytes to macrophages. 147,148 Only a few additional details of some of the key processes are provided here.

As noted before, initial tethering and rolling of leukocytes are mediated, in part, by P-selectin and E-selectin on activated endothelial cells and L-selectin on leukocytes. PSGL1 (P-selectin glycoprotein ligand 1) can bind all these selectins. Other ligands can also bind E-selectin. Nonactivated integrins can bind weakly to VCAM-1 and ICAM-1 and can also mediate rolling. However, activated integrins are generally considered more important in leukocyte firm adhesion and spreading. P-selectin is also expressed on activated platelets. P-selectin is presynthesized and stored in Weibel-Palade bodies and can be expressed on the cell surface rapidly by fusion of the Weibel-Palade body with the cell membrane. In contrast, E-selectin is not stored and must be synthesized de novo; thus, levels rise more slowly on activation.

The early recognition of the predominance of macrophages in human and experimental plaques focused most attention on the early entrance of these cells into the intimate space. However, the importance of other cells has been increasingly recognized. Further stimulation of macrophages or dendritic cells by Th1 cells as well as mutual stimulation by macrophages and dendritic cells seems to establish inflammatory cells within the intima as long as any inciting risk factors (such as hyperlipidemia) are present. Neutrophils may become particularly important in destabilizing advanced lesions. ^{140,143}

In a startling set of experiments, an extensive network of dendritic cells were found to be the earliest inhabitants of this space, concentrated clearly in atherosclerosis-prone areas of disturbed flow. This network was established in all specimens examined even in infancy or early childhood in the absence of any risk factors, well before the presence of any evidence of macrophage accumulation or lipid-filled cells, and was universally present in normolipidemic animals as well. 149 The participation of these cells in addition to early appearance of proinflammatory Th1 lymphocytes, which may respond to the self antigen HSP-60, together with the observation that patients with CAD were found to have increased plasma titers of anti-HSP-60 antibodies that strongly cross-reacted to certain Cytomegalovirus antigens has raised considerable interest. 150 Evidence has been presented that immunization of LDLR knockout mice with the bacterial homologue HSP-65, HSP-60, or desensitizing peptide fragments shifts lymphocyte counts to greater numbers of regulatory Th2 (anti-inflammatory) in relation to Th1 cells with increased IL-10 production and substantially reduces atherosclerosis by 50% to as much as 80%. 151,152 Whether such desensitization therapy may be effective to prevent progression of human atherosclerosis is unknown.

Because the movement of leukocytes has been so extensively studied in the context of many diseases, the elegant mechanisms controlling their movement and chemotaxis will not be discussed in detail. It is recommended that the interested reader consult recent textbooks for the elegant coordination of these events, in which GTPase molecules Cdc42 and Rac serve as both controlling "switches" and membrane attach ment points for growing and branching actin microfibers that provide a pushing force to propel the cell forward, while Rho works at the back of the cell to promote uropod retraction. Kinesin motors serve to recycle integrins and other key cell machinery forward while other signaling mechanisms maintain cell polarity.

8

Signaling	ytokines
genes	Effect
Secreted by <i>Macrophages</i> in response to Toll Recept Receptor Binding	or and Scavenger
IL-1 a	you
IL-1 0	4
IL-6	NS
gp130 (IL6ST, in humans)	4
IL-12	44
IL-18	4
TNF- α	44
MIF	44
IL-2	4
IEN F	1 to 11
IFN- Γ Secreted primarily by <i>Th2 cells</i> after activation by An (Primarily Anti-Inflammatory) IL-4	
Secreted primarily by <i>Th2 cells</i> after activation by An (Primarily Anti-Inflammatory) IL-4	
Secreted primarily by <i>Th2 cells</i> after activation by An (Primarily Anti-Inflammatory)	tigen Presentation
Secreted primarily by <i>Th2 cells</i> after activation by An (Primarily Anti-Inflammatory) IL-4 IL-5 IL-10	tigen Presentation NS 4
Secreted primarily by <i>Th2 cells</i> after activation by An (Primarily Anti-Inflammatory) IL-4 IL-5 IL-10	tigen Presentation NS 4
Secreted primarily by <i>Th2 cells</i> after activation by An (Primarily Anti-Inflammatory) IL-4 IL-5 IL-10 Secreted primarily by <i>T Regulatory Cells</i> TGF- p Secreted by <i>Mast Cells</i>	tigen Presentation NS 4
Secreted primarily by <i>Th2 cells</i> after activation by An (Primarily Anti-Inflammatory) IL-4 IL-5 IL-10 Secreted primarily by <i>T Regulatory Cells</i> TGF- p	tigen Presentation NS 4
Secreted primarily by <i>Th2 cells</i> after activation by An (Primarily Anti-Inflammatory) IL-4 IL-5 IL-10 Secreted primarily by <i>T Regulatory Cells</i> TGF- p Secreted by <i>Mast Cells</i>	tigen Presentation NS 4 4
Secreted primarily by <i>Th2 cells</i> after activation by An (Primarily Anti-Inflammatory) IL-4 IL-5 IL-10 Secreted primarily by <i>T Regulatory Cells</i> TGF- p Secreted by <i>Mast Cells</i> GM-CSF Miscellaneous	tigen Presentation NS 4 4 4 4 to 4
Secreted primarily by <i>Th2 cells</i> after activation by An (Primarily Anti-Inflammatory) L-4 L-5 L-10 Secreted primarily by <i>T Regulatory Cells</i> TGF- p Secreted by <i>Mast Cells</i> GM-CSF Miscellaneous L-1Ra (TG)	tigen Presentation NS 4 4 4 4 to 4

NS, not significant; TG, transgenic with overexpression.

Once early inflammatory changes have been established, particularly with recruitment of the various leukocytes to the subendothelial space, numerous cytokines are secreted by activated cells that reinforce the inflammatory response and further activate endothelial cells as well as surrounding leukocytes. This exchange of cytokines acts as a positive feed back loop to ensure a vigorous response to perceived threats. In Table 8-3, a list is provided of some of these cytokines as well as other genes affecting early stages of endothelial activation and inflammation and effects on atherosclerosis of genetic manipulation. The receptors and intracellular signal ing in response to these cytokines have been the subject of much study, and many illustrations of these pathways are available for free download from commercial sources on the Internet (eg, http://www.sabiosciences.com/pathwaycentral and http://www.cellsignal.com/reference/pathway/ index.html) as well as a recent on-line textbook of cell signal ing pathways (http://www.cellsignallingbiology.org/). They will not be discussed further here.

While reviewing Tables 8-1 through 8-3, the reader should keep in mind that the apparent impact on atherosclerosis of manipulating a given gene can change over time. Most of these models have double knockouts; the first knockout (of the apo E

gene or the LDL receptor) results in hyperlipidemia, and the second knockout is of the test gene under study. Short-term studies generally report a greater percentage reduction in atherosclerosis due to gene knockout tests than longer-term studies do. For example, in one study, at 5 weeks there was a clear effect of even heterozygous CCR2 (the receptor for MCP-1) knockout, with atherosclerosis clearly increasing progressively from CCR2 / (strongly protected) to CCR2 / to CCR2 / (wild type). By 9 weeks, however, the heterozygous knockout had caught up to the wild type. By 13 weeks, even the homozygous knockout mice had atherosclerosis extent similar to that for wild type at 9 weeks, although the trajectories for atherosclerosis remained different between CCR2 / and CCR2 / little. ¹⁵³

What do such observations imply for gene effects in human atherosclerosis, a disease that develops over decades? Even with powerful (but presumably rare) effects on a clearly important gene, there may be no observable effect in the long run in a redundant system. This principle may help explain the extreme difficulty that human gene hunters have had in identifying consistently replicable genetic associations with atherosclerosis, particularly involving initiating pathways (endothelial activation and inflammation). A corollary may be that interventions that have global impact on such pathways (such as

control of hyperlipidemia or a reduction in blood pressure) will likely have more success than efforts to affect a single genetic element. An additional consideration with regard to inflammatory pathways is that a focused intervention such as that noted for CCR2 knockout that resulted in marked reductions in monocytes may be associated with unacceptable increases in susceptibility to infection.

Effects of Selected Risk Factors on Initiation of Atherosclerosis

Blood Pressure

Pressure, stretch, and flow are the first risk factors to which the endothelium is exposed. Blood pressure in the arterial range is an essential requirement for the development of atherosclerosis. Venous atherosclerosis does not occur even in patients with homozygous familial hypercholesterolemia. ¹⁵⁴ The relatively rapid progression of atherosclerosis in saphenous veins used in coronary artery bypass suggests that there is nothing uniquely resistant about veins themselves. Normally, the pulmonary arterial circulation with its systolic pressures of 12 to 22 mm Hg is another entirely protected site. Nevertheless, with pulmonary hypertension, atherosclerotic plaques are commonly seen. ¹⁵⁵

The major mechanism whereby pressure promotes - atherosclerosis appears to be through increased pressure-driven convection of LDL and other lipoproteins into the intima. ¹⁵⁶ For example, LDL accumulation in the intima of pressurized rabbit aorta increased 44-fold when pressure was raised from 70 to 160 mm Hg. ¹⁵⁷ Stretch may also play a role. ¹⁵⁸ Approximately 90% of the convection of LDL into the subendothelial space appears to be through gaps created between mitotic endothelial cells in areas of low shear stress, whereas only 10% was estimated to enter by way of transcellular vesicular traffic. ¹⁵⁹ A thinner glycocalyx, also associated with atherosclerosis-prone areas with low shear stress, also appears to contribute to greater LDL permeability. ¹⁶⁰

Exercise

Even modest exercise serves as a potent stimulus to improve endothelial function as measured by vasodilation

Stimulation of flow has a number of additional effects. An acute increase in flow with exercise results in a transient increase in endothelial superoxide anion generation by mitochondria and NADPH oxidase. 165 This superoxide is rapidly converted to hydrogen peroxide. The modest, controlled burst of hydrogen peroxide, in turn, also acts to promote increased expression and activity of eNOS, possibly (as noted before) through altered expression of NF- K B as well as stimulation of HSP-90 (which strongly supports eNOS activity) through stimulation of HNF1. Importantly, when high levels of human catalase were expressed in endothelial cells of exercising mice, blocking signaling by hydrogen peroxide, the expected increase in eNOS expression in response to exercise was entirely blocked. 166

Endothelial cells also adapt to chronic laminar flow by stimulation of a number of antioxidant defenses (at least in part through Nrf2 signaling), resulting in increased expression of extracellular superoxide dismutase and Cu/Zn SOD as well as a decrease in NADPH oxidase. 165 Similar adaptive changes occur with exercise training, explaining the longer term antioxidant effects of exercise despite acute modest stimulation of superoxide and hydrogen peroxide production during acute bouts of exercise. 161,166 Parallel benefits from exercise on insulin signaling pathways in skeletal muscle appear to be blocked by large doses of oral antioxidants, suggesting potential deleterious effects of such interventions. 167 Finally, exercise was shown to clearly reduce atherosclerosis in apo E-deficient mice together with marked reduction of macrophage and Th1 cell accumulation in the intima. This effect of exercise was completely blocked by inhibition of eNOS. 164 Other effects of exercise on endothelial function, including increased PGI 2, thrombomodulin, and plasmin, have also been reviewed. 168

Hyperlipidemia

expression. 164

Extensive effects of hyperlipidemia to initiate atherosclerosis have been reviewed earlier. Not only does hyperlipidemia activate the endothelium and otherwise promote atherosclerosis, but effects on circulating white cells are also apparent. Thus, expression of CCR2, the cognate receptor on monocytes for MCP-1, was approximately twofold higher hypercholesterolemic patients with average LDL of 167 mg/dL compared with persons with LDL of 80 mg/dL. Incubation of monocytes in hypercholesterolemic serum, without LDL oxidation, led to similar marked increases of CCR2, apparently through uptake by the LDL receptor. Additional unexpected effects of hypercholesterolemia include suppression of the inward rectifying potassium current in endothelial cells (which might be expected to lead to diminished NO response to acute changes in flow). 169 Hypercholesterolemia affects phosphate dylinositol 4,5-bisphosphate (PIP 2)-sensing amino acids in inwardly rectifying K (Kir2) channels, not by altering membrane physical properties. 170 Interestingly, patients with familial hypercholesterolemia were shown to have reduced glycocalyx volume, which could be partially restored by treatment with

rosuvastatin. 171

Hypercholesterolemia and particularly hypertriglyceridemia appear to increase the activity of xanthine oxidoreductase (XOR). 172 There may be both greater conversion in the liver of the enzyme from the dehydrogenase form to the oxidase form (accompanied by greater production of superoxide) and increased release of the oxidase from the liver. Xanthine oxidase is then thought to adhere to the glycocalyx of endothelial cells, where it can contribute to endothelial oxidative stress and promote endothelial dysfunction that may be improved by XOR inhibitors such as allopurinol. 173 XOR knock out studies are not 8 feasible in mice as the pups are stunted and live only 4 to 6 weeks, presumably because of renal failure. 174 Nevertheless, tungsten administration (a relatively specific inhibitor of XOR) to apo Edeficient mice fed a Western-type diet for 6 months resulted in 83% reduction of aortic atherosclerosis. 175

HDL Effects

Besides reverse cholesterol transport, HDL has been shown to mediate a number of protective effects on the endothelium. HDL protects endothelial cells from apoptosis induced by oxLDL, TGRL, TNF- a , and activated complement. HDL stimulates NO production by eNOS and increases prostacyclin (PGI 2) n> production through COX-2. Finally, HDL suppresses the expression of VCAM-1, ICAM-1, and E-selectin by endothelial cells as well as other signs of endothelial activation after exposure to oxLDL or cytokines. 176 These effects are mediated by both apo AI and lipid components carried in HDL, particularly sphingosine 1-phosphate (S1P) and related lysosphingolipids.

There are five known S1P receptors, S1P1 through 5. These are G protein-coupled receptors (GPCR) with the potential to be coupled to a variety of heterotrimeric G proteins with diverse and frequently divergent functions. The atheroprotective functions of HDL on endothelial cells appear to be mediated by S1P signaling primarily through the S1P1 receptor and to a lesser extent through S1P3. 177,178 HDL carries more than 50% of the S1P in plasma. 179,180 Activation of the S1P1 receptor recapitulates a number of the protective features of prolonged exposure to laminar flow (see Fig. 8-7). These effects appear to be mediated by ERK1/2, PI3K, Akt, and Rac1 activation, with resulting increased barrier integrity , increased NO production, suppression of inflammatory responses, promotion of cellular antioxidants, and increased cell survival (by suppression of caspases). Activation of these pathways is supported by and possibly dependent on the binding of apo AI to endothelial SR-B1. Binding to SR-B1 not only brings the HDL into proximity with S1P1 receptors but also supports cytosolic signaling to activate Src. Src can then activate MEK1/2 followed by ERK1/2 as well as PI3K. PDZK1 is an adapter or scaffolding protein that binds to the cytosolic side of SR-B1. PDZK1 is required for Src activation by HDL. Indeed, PDZK1 knockout completely blocked HDL-mediated increases in eNOS and promotion of endothelial repair after injury. 181 HDL may also inhibit monocyte activation after interactions with T cells. 177

Another mechanism by which HDL protects endothelial cells and promotes NO production is by removal of the oxy sterol 7keto cholesterol by way of the ABCG1 transporter. 182 This oxidized sterol built up when mice were fed a Western type diet. HDL also helps prevent LDL oxidation or detoxifies oxidized phospholipids in LDL through the HDL-associated enzyme PON1. PON1 knockout mice express increased atherosclerosis, whereas overexpression of PON1 was protective. 183 These findings contrast with the much more variable outcomes associated with milder effects on PON1 caused by common human variants. 184



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Whereas other risk factors may recapitulate a form of endothelial activation that in some settings would be advantageous (such as in infection or physical injury), the endothelial response in diabetes is truly maladaptive and finds few if any parallels in normal physiology. Therefore, the term *endothelial dysfunction* applies most aptly in diabetes. Unlike other risk factors, hyperglycemia leads to endothelial dysfunction that is systemic, resulting in both aggravation of atherosclerosis and a microvascular disease that is unique to diabetes. In type 2 diabetes, excess free fatty acid exposure and insulin resistance appear to further exacerbate this response.

An elegant, unified model to explain the seemingly disparate aspects of endothelial dysfunction in diabetes has been forwarded. 185,186 The endothelial cell is one of a handful of cell types that are unable to regulate the entry of glucose into their cytoplasm. Thus, hyperglycemia leads to unregulated uptake of glucose through GLUT1 transporters and elevated intracellular glucose in endothelial cells. Rapid flux of substrates through the glycolytic pathway then ensues, leading to an abundance of pyruvate for transfer into mitochondria, synthesis of acetyl coenzyme A, and generation of reducing equivalents (NADH, FADH 2) by the citric acid cycle. The transfer of high-energy electrons from NADH and FADH 2 to complexes of the electron transport chain provides energy to pump hydrogen ions from the matrix into the mitochondrial intermembrane space, thereby generating the hydrogen ion gradient that subsequently drives ATP synthesis through F₀F₁ complexes. With an overabundance of energy substrate, the hydrogen gradient increases to a critical level, at which point the final electron transport to complex IV is impaired (where water is normally formed from controlled electron transport to oxygen). The "backed up" electrons then begin to be transported from coenzyme Q 10 (ubiquinone) to oxygen directly, forming superoxide anion. (Paradoxically, a similar phenomenon occurs with hypoxia.) This formation of superoxide anion can be blocked by dissipating the hydrogen ion gradient (eg, by overexpression of uncoupling proteins) or by preventing the transfer of electrons into the transport chain. In larger arteries, uncontrolled endothelial uptake and oxidation of free fatty acids, which are excessively abundant in type 2 diabetes, further exacerbate the energy surfeit and excess generation of superoxide anion.

Although superoxide anion does not penetrate mem branes, a substantial amount of the excess superoxide is formed on the outer side of the mitochondrial inner mem brane and can be transferred to the cytosol through the voltage-dependent anion channel in the outer mitochondrial membrane. 58 A modest or transient increase in superoxide production might be expected to produce primarily a transient rise in hydrogen peroxide and potentially stimulate an antioxidant response through Nrf2 signaling, as apparently occurs with exercise. Such a response is seen with short-term incubation of endothelial cells in high glucose. However, prolonged incubation, as in diabetes or with frequently repeated although transient exposures to high glucose, appears to result in sufficient superoxide production to over whelm SOD antioxidant defenses and results in nuclear (as well as mitochondrial) DNA damage, such as single-strand breaks. In response, a DNA repair enzyme, poly(ADP-ribose) polymerase 1 (PARP-1), is activated. PARP-1 uses NAD as a substrate and adds long ADP-ribose chains to various proteins and transcription factors, to itself, and, importantly, to histones. ADP-ribosylation of histones alters their conformation and allows greater access to the damaged DNA for other repair enzymes. Hydrogen peroxide, although less reactive than other ROS, can also activate PARP-1 (perhaps through formation of highly reactive hydroxyl radical), but to a lesser extent than superoxide and peroxynitrite. 187 PARP-1 can signal for apoptosis or extensive inflammation in the setting of extensive DNA repair, and pharmacological inhibition or genetic deletion of PARP-1 decreased inflammation and reduced atherosclerosis in hyperlipidemic mice. 188 Activated PARP-1 leads to inhibition of glyceraldehyde-3-phosphate dehydrogenase with accumulation of the 3-carbon substrates, with subsequent promotion of AGE synthesis as well as diacylglycerol production with stimulation of PKC

Stimulation of the expression of Nrf2 (by inactivation of Keap1 with sulforaphane, a naturally occurring substance in broccoli) upregulated antioxidant defenses and led to a marked reduction in the adverse effects of hyperglycemia, further supporting the role of ROS in mediating endothelial dysfunction in diabetes. ¹⁸⁹ Lipoic acid is an important mitochondrial antioxidant, and supplementation was shown to decrease atherosclerosis in hyperlipidemic, diabetic mice. ¹⁹⁰ Heterozygous deficiency of lipoic acid synthase in such mice resulted in a 48% increase in atherosclerosis extent. ¹⁹¹

There is growing evidence for selective endothelial insulin resistance for the vasodilating Akt pathway while the vasoconstricting ERK1/2 pathway (through Shc, Grb2-Sos, Ras, Raf, and MEK1/2) is left intact. The vasoconstriction seems to be particularly promoted through the activation of PKC O by fatty acids, such as palmitate. PKC O both inhibits insulin signaling through IRS-1 and Akt and promotes signaling through the ERK1/2 pathway. 192,193 Perhaps more relevant are increases in adhesion molecule expression in endothelial cells after exposure to insulin. Insulin-induced expression of VCAM-1 in endothelial cells was recently shown to be mediated through the insulin-like growth factor 1 receptor and could be completely abrogated by inhibiting the MAPK p38. Interestingly, insulin also stimulated the expression of ICAM-1 by way of the insulin receptor. In both cases, MEK1 appeared to be involved but primarily by activating p38 rather than ERK1/2. 194 Endothelial dysfunction (including impaired eNOS phosphorylation and production of NO) was clearly induced recently by exposure to saturated free fatty acid (particularly palmitate) regardless of insulin signaling (with use of mice lacking insulin receptors in all the vasculature or knockout of Akt). 195 These mice also developed hypertension.

Autoimmune and Inflammatory Disease

Increased risk of atherosclerotic disease has been documented for several autoimmune diseases, including systemic lupus erythematosus, 196-201 rheumatoid arthritis or elevated rheumatoid factor, 202,203 systemic sclerosis, 204 and psoriasis. 205-²⁰⁷ It would be reasonable to expect that the elevated plasma cytokines, including TNF- a and IL-6, 208,209 seen particularly in systemic lupus erythematosus, would activate endothelial cells and thereby initiate atherosclerotic processes, particularly at atherosclerosis-prone sites. Other mechanisms have recently been forwarded. Mice that lack functional Fas ligand or Fas have impaired ability to clear apoptotic debris and have features of autoimmune disease. When crossed with apo E-deficient mice, these mice have increased atherosclerosis. 210,211 Apoptotic debris may present autoantigens to dendritic cells and thereby lead to autoantibody production and increased circulation of immune complexes, which could cause immune injury and promote inflammation. Alternatively, apoptotic debris itself could directly promote inflammation in atherosclerotic plaques.

Several older studies illustrated a potential synergism between immune injury and hyperlipidemia. Rabbits made only mildly hypercholesterolemic (200 to 250 mg/dL) and subjected to repeated injection of horse serum (a form of immunologic assault with antigen-antibody deposition on lesion-prone areas of endothelium) developed atherosclerosis that was more similar to human atherosclerosis than the usual severe hypercholesterolemia-induced lesions in rabbits. The rabbits appeared to be protected by antihistamines in this

setting. ²¹² A counterpart in humans was suggested to be the finding of increased levels of serum antibodies to heat-dried cow's milk and boiled egg in patients with coronary heart disease. ²¹³ The display of HSP-60 on endothelial cells in association with activation of endothelial cells from a variety of initiating factors could lead to autoimmune reactions, endothelial damage, and increased atherosclerosis, especially coupled with hyperlipidemia and other risk factors. ⁵²

A large literature has accumulated relating to immune activation by repeated or latent infections, which suggests another potential mechanism for endothelial activation and atherosclerosis initiation. ²¹⁴ Macrophage activation may be involved as well. Infection of apo E-deficient mice with *Chla-mydia pneumoniae* ²¹⁵ or herpesvirus ²¹⁶ accelerated -atherosclerosis. Indeed, so did nonspecific activation of TLR2 receptors. ¹¹² However, treatment of coronary patients with macrolide antibiotics failed to alter subsequent event rates. ²¹⁷⁻²¹⁹ Thus, causality for the infectious disease hypothesis remains to be proven. Furthermore, if the relationship to infectious disease requires only exposure resulting in an immune response, not continued infection, then the hypothesis may not be proven by any antimicrobial intervention.

Cigarette Smoking

Smoking has long been considered an endothelial cell stressor and activator. A genome-wide analysis of expression changes in 35,000 genes was recently reported after exposure of cultured endothelial cells to cigarette smoke extract. 220 There was massive upregulation of genes related to the unfolded protein response, thought to be due to immense oxidative stress. Generation of free radicals was apparently mediated by metals as well as by reactive species in the extract. This was accompanied by mitochondrial dysfunction, upregulation of HSF1 and heat shock proteins (including HSP-60), and cell cycle arrest. These observations are consistent with prior reports of the pro-oxidant effects of smoking leading to activation of endothelial cells. ²²¹ The rapid reversibility of cardio vascular risk after smoking cessation suggests that this activation affects not only early endothelial activation but precipitating factors such as clotting and stability of advanced plaques as well (probably also through inflammatory effects). 222 Nevertheless, some evidence suggests that persistent inflammatory effects can last for years after smoking cessation. 223

PROMOTION OF ATHEROSCLEROSIS

The promotion of atherosclerosis centers on the development of foam cells, their retention in or egress from the intima, their apoptosis or necrosis, and formation of the necrotic core. Insudation of lipoproteins into the subendothelial space and lipoprotein retention and modification are thought to be the major drivers of this stage of atherogenesis. The risk of coronary events is clearly proportional to the exponent of LDL plasma concentrations (hence the frequently replicated log-linear relationship, with a 30% increase in risk associated with a 30 mg/dL linear rise in plasma LDL). ²²⁴ Other apo B-containing lipoproteins are also proatherogenic; the order of atherogenicity, as demonstrated by matching cholesterol levels in apo E and LDLR knockout mice, ²²⁵ is roughly as follows: smaller LDL > larger LDL > \$\bar{\scrt{S}}\$ -VLDL > IDL > smaller VLDL > larger VLDL.

These findings are consistent with findings in humans. Patients with homozygous familial hypercholesterolemia (LDLR mutations) have the highest risk for CAD, followed by LDLR heterozygotes, type III patients, then type IV and possibly type V patients. Interestingly, human subjects with heterozygous LDLR mutations and either heterozygous or homozygous for apo E2-2 (and who clearly manifest type III hyperlipidemia in addition to high LDL) do not have a higher risk than that of other persons with familial hypercholesterolemia. ²²⁶⁻²²⁸ This may be due to a

reduction in conversion of remnants to LDL and somewhat reduced LDL levels in this combined lipid disorder. In contrast, mice with combined LDLR and apo E deficiency suffer extremely accelerated atherosclerosis and experience spontaneous myocardial infarction, which also seems to be related to endothelin signaling as a precipitating event. ²²⁹ Low levels of HDL are clearly associated with elevated atherosclerosis risk. The association of low HDL with elevated triglycerides can be associated with remarkably elevated risk. ^{230,231} The remainder of this section focuses on mechanisms whereby lipoproteins promote foam cell formation. Genetic determinants of lipo - 8 protein levels are treated elsewhere in this volume.

As reviewed earlier, endothelial cells are activated by several different forms of modified lipoproteins. Macrophages share with endothelial cells many common receptors and activating mechanisms. In general, macrophages will be activated once they reach the subendothelial space through signaling with activated endothelial cells. However, they become more fully activated, expressing various scavenger receptors with increased secretion of various cytokines as well as activation of myeloperoxidase, after they encounter proinflammatory, modified lipoproteins. These changes generally occur concomitantly during foam cell formation.

Lipoprotein Retention and Modification

The level of LDL found in the interstitial fluid of the intima is approximately double the plasma concentration. ²³² As demonstrated recently in human autopsy specimens, lipoprotein entry and retention in the intima precedes macrophage infiltration into this space. ²³³ Once partially activated - macrophages enter the subendothelial space, they encounter trapped lipoproteins. As shown by the classic experiments of Brown and Goldstein, unmodified LDL cannot lead to foam cell formation as the uptake of native, unmodified LDL is by way of the LDLR, which is downregulated once cellular cholesterol - stores are replete. ²³⁴ Ş -VLDL was the only native lipoprotein that was taken up into macrophages and that led to foam cell formation without modification in those pioneering studies. These earlier observations have been amply verified. ²³⁵

A number of LDL modifications have now been shown to also be sufficient in the formation of foam cells. Many of these modifications are not by way of oxidation. ²³⁶ Clearly, the LDL and other lipoproteins must remain trapped in the intima long enough for modification to occur and for macrophages to take them up for atherosclerosis to progress. The importance of retention of LDL by binding to proteoglycans in the intimal extracellular matrix (particularly to those bearing chondroitin sulfate) was clearly demonstrated by the decrease in atherosclerosis seen in mice whose apo B lacked a proteoglycan binding determinant. ²³⁷ Other proteoglycan binding sequences appear to be present in apo B48 as well. 8 Lipoprotein lipase (LPL), when expressed by macrophages, can provide a nonenzymatic bridge between lipoproteins and proteoglycans and promote retention in the arterial wall and increased atherosclerosis (as shown by bone marrow transplantation studies with LPL-deficient macrophages). ²³⁸ Conversely, when LPL is playing its more commonly considered role on endothelial cells, it promotes lipoprotein delipidation of TGRL and is protective, especially when it is overexpressed in LDLR knockout mice. 239

One of the most important mechanisms for both retention and modification of trapped lipoproteins appears to be mediated by secreted phospholipases (sPLA $_{\rm 2}$), particularly groups III and V sPLA $_{\rm 2}$. $^{\rm 240}$ Apo E knockout mice fed an atherogenic diet and overexpressing human group III sPLA $_{\rm 2}$ had a more



than twofold increase in atherosclerosis. 240 LDL modified by group V sPLA $_2$ aggregates and binds proteoglycans more avidly and is taken up into macrophages, leading to foam cell formation. Accelerated atherosclerosis was seen in LDLR knockout mice when group V sPLA $_2$ was overexpressed; reduced atherosclerosis was seen in mice with bone marrow transplants from mice deficient in group V sPLA $_2$. 241 Group IIa sPLA $_2$ may also contribute but is less active than the group V enzyme. 242 Group V sPLA $_2$ is the major sPLA $_2$ enzyme released by activated endothelial cells. 243,244

Sphingomyelinase is another enzyme released by activated endothelial cells and other lesional cells that has been implicated in LDL modification. Importantly, sphingomyelin produces ceramides in the lipoprotein and promotes greater retention by proteoglycans and greatly promotes uptake and foam cell formation for several lipoprotein types. Deficiency of acid sphingomyelinase in LDLR knockout mice resulted in a 55% decrease in atherosclerosis. ²⁴⁵ Not only might sphingomyelinase be important within the subendothelial space, but a fraction of plasma LDL appears to carry substantial sphingomyelinase activity. These LDL were also electronegative. Electronegative LDL can also carry substantial PAF-AH (also referred to as Lp-PLA 2). Electronegative LDL, when incubated in vitro, were shown to degrade their own phospholipids in vitro, rendering them highly subject to aggregation and macrophage uptake. This activity was not blocked by antioxidants and could represent an important means of LDL modification. However, the source of the activity was apparently not secreted sphingomyelinase. 125 It would be of great interest to assess the effect of knocking out both group V sPLA 2 and sphingomyelinase in hyperlipidemic animals as these two enzymes appear to work synergistically to modify LDL.

Much has been written about oxidative modification of lipoproteins. Certainly, when oxidized by various methods, including incubation with cultured endothelial cells (which are generally at least partially activated in most in vitro settings), LDL and other lipoproteins are taken up avidly by macrophages with resultant macrophage activation, proliferation, and foam cell formation. 97 Some of these effects are likely to be mediated by oxidation of phospholipids ²⁴⁶; oxidized cholesteryl esters ²⁴⁷ and changes to apo B characterized by chlorination of tyrosine and nitrations by myeloperoxi dase are important as well. 101 Furthermore, oxidation of HDL clearly impairs antiatherosclerotic activity. 101 These and many other observations make a persuasive argument that oxidation of lipoproteins contributes to atherosclerosis. 97,248 However, human clinical trials with antioxidants have been resoundingly negative (particularly the larger and more recent ones).

A potential explanation to resolve this apparent discrepancy may be, once again, redundancy. Here, the redundancy involves multiple possible lipoprotein modifications that could all lead to foam cell formation. One additional consideration may be chemical oxidative The footprint predominantly on LDL from human atherosclerotic plaque clearly points to myeloperoxidase as the source of pro-oxidant activity. 249 This would suggest that activated macrophages and possibly (in more advanced lesions) neutrophils are the main cause of such oxidized LDL. This brings into question the primacy of oxLDL (or possibly even mmLDL) in foam cell formation because activated macrophages would have to already be present in the lesion to initiate such oxidation . Whereas oxLDL generated by myeloperoxidase would undoubtedly be atherogenic and promote further foam cell formation as well as inflammation and apoptosis, it would seem unlikely to be an initial cause of macrophage activation (and perhaps not even a necessary one). Furthermore, the LDL could be oxidized after it was taken up into the macrophage.

Unfortunately, standard mouse models of atherosclerosis are not of much help in resolving this issue as mice express much lower levels of myeloperoxidase in their macrophages than is seen in human lesions. Furthermore, knockout of mouse myeloperoxidase does not decrease atherosclerosis. However, when human myeloperoxidase is expressed in the mouse macrophages, which results in increased levels, atherosclerosis is increased. ¹⁰¹ These considerations support the notion that myeloperoxidase is potentially proatherogenic but redundant, as mouse atherosclerosis can progress extensively in its absence.

Carbamylation is a recently recognized form of LDL modification that may be particularly important in renal disease and smoking. A small fraction of urea can spontaneously and reversibly form highly reactive cyanate, which can then bind to lysine residues in proteins, forming homocitrulline. Many proteins in uremia have been noted to be altered, but carbamylated LDL from uremic patients or prepared in vitro activated endothelial cells primarily by way of LOX-1 on endothelial cells. ²⁵⁰ Furthermore, carbamylated LDL binds to macrophage SR-AI receptor and causes foam cell formation. 251 Importantly, thiocyanate, which is markedly increased in the plasma of cigarette smokers, can be converted to cyanate by myeloperoxidase, which might mediate a substantial portion of carbamylation in vivo. Protein homocitrulline levels were found to be strongly related to cardiovascular risk with an odds ratio of 7 to 8 in the top versus bottom quartile. ²⁵¹

Uptake of Modified Lipoproteins to Form Foam Cells

Traditionally, scavenger receptors have been considered the primary mechanism whereby modified lipoproteins are taken up relentlessly by macrophages, leading to foam cell formation and promotion of atherosclerosis. However, recent evidence points to non-scavenger receptor uptake as a quantitatively more important pathway. LDL treated with group V sPLA 2 is taken up by macropinocytosis, which is mediated by syndecan 4. This process is accelerated by macrophage activation and can lead to foam cell formation. Unlike uptake of oxLDL through scavenger receptors, macropinocytosis is blocked by inhibition of PI3K. ²⁵² Thus, foam cells can clearly be formed independently of scavenger recept tors. In a sense, then, scavenger receptors and uptake of oxidized LDL may be considered yet another example of redundant pathways.

Oxidatively modified LDL as well as some other modified LDL and \$ -VLDL are taken up by a variety of scavenger receptors. ²⁵³ These include the class A (AI, AII, MARCO, SRCL, SCARA5), class B (CD36, SR-B1), class C (dSR-C1, in *Dro sophila* only), class D (CD68, an endosomal receptor), class E (LOX-1), class F (SREC-I, SREC-II), class G (SR-PSOX = CXCL16), class H (FEEL-1, FEEL-2, primarily bacterial binding), and class I (CD163, may help clear hemoglobin) receptors. A number of these receptors bind acetylated LDL and especially oxLDL (SR-AI/II, MARCO, SRCL, SR-B1, LOX-1, SREC-I, FEEL-1/2). CD36 also binds VLDL and TGRL remnants. However, only SR-A and CD36 have been subject to extensive studies.

Importantly, all these scavenger receptors bind alternative ligands, which may reflect their physiological roles. Thus, scavenger receptors bind various bacteria, bacterial lipopolysaccharide, lipoteichoic acid, microbial diglycerides, apoptotic debris, amyloid, and AGEs. Macrophages from mice deficient in SR-AI/II bound AGEs much less avidly. Also, the mice were more prone to death from infection with *Listeria* and particularly with HSV-1 infections. ²⁵⁴ Thus, macrophage scavenger receptors play a major role in recognizing PAMPs and hence participating in innate immunity. Later it was

streptococcus and Streptococcus pyogenes but could also bind Neisseria meningitidis, Staphylococcus aureus, and Escherichia coli. ²⁵³ Binding of CD36 can trigger inflammatory activation of NF-KB and MAPK cascade signaling by interactions with Src kinases and tolllike receptors or certain integrins as coreceptors with CD36. ²⁵⁵ Nevertheless, scavenger receptors can also play an important antiinflammatory role in clearing apoptotic debris.

by the MSR gene) and CD36 showed no alteration in the extent of atherosclerotic lesions in apo E knockout mice. 256 Importantly, however, these MSR/CD36/apo E triple knockout mice did have much less apoptosis and necrosis and fewer inflammatory markers in their lesions. The transcription factor STAT1 was also recently shown to be critical for apoptosis in this setting. ²⁵⁷ Thus, scavenger receptors may be more important for later progression of atherosclerosis, including formation of the necrotic core and plaque instability.

Cholesterol Homeostasis in Macrophages

As macrophages indulge, atherosclerotic lesions bulge. ²⁵⁸

As macrophages take up modified LDL and TGRL remnants, cholesteryl ester in lysosomes is hydrolyzed by acid choles teryl ester hydrolase (CEH). The released free cholesterol is then esterified by ACAT1 and stored in intracellular lipid droplets. These droplets are surrounded by several proteins including perilipin, ADRP (adipose differentiation-related protein), TIP47 (tail-interacting protein of 47 kDa), and S3-12. ²⁵⁹ Cholesteryl ester stored in these droplets can be hydrolyzed by neutral CEH. Free cholesterol released in this manner appears to be targeted for efflux from the macrophage through the ABCA1 receptor, which interacts primarily with lipid-free apo AI or very small pre- \$ HDL; the AGCG1 or AGCG4 receptors, which release cholesterol to larger HDL; or the SR-B1, also to HDL. Macrophages also constitutively secrete apo E, which can act, with phospholipid, as a free cholesterol acceptor, forming discs, much like nascent HDL. ²⁶⁰

Some intracellular free cholesterol is spontaneously or enzymatically (by CYP27A) oxidized to oxysterols (such as 25- or 27-hydroxycholesterol), which bind to and activate the liver X receptor (LXR). When thus bound, LXR dimerizes with the retinol X receptor (RXR) and affects transcription of genes involved with cellular cholesterol balance. Thus, activated LXR increases transcription of Niemann-Pick type C 1 and 2 proteins (which are involved in intracellular cholesterol trafficking), ABCA1, ABCG1, apo E, and PLTP (phospho lipid transfer protein). Activated LXR can also be sumoylated and, as such, strongly suppresses transcription of a number of proinflammatory Pharmacological activation of LXR inhibits atherosclerosis, whereas knockout results in increased atherosclerosis. 261,262 Activation of PPARy also upregulates LXR expression but induces CD36 expression as well. ²⁶³ Interestingly, the fatty acid-binding protein aP2 appears to bind natural ligands of PPAR y, and knockout of aP2 leads to greater stimulation of PPAR y and suppression of atherosclerosis. ²⁶⁴

In considering the effects of various genes on cholesterol homeostasis in macrophages, one key fact must be recognized . Excess free or unesterified cholesterol (FC) is toxic to cells. 15,265 Foam cells can accumulate large amounts of cholesteryl ester with relative impunity. However, in the process of cholesteryl ester hydrolysis in lysosomes, free cholesterol can rise to high levels. This is the location of most free cholesterol in foam cells found in advanced plaques; yet it is excess FC transported to endoplasmic reticulum that may lead to the earliest cellular dysfunction, possibly by precipitating release of endoplasmic reticulum calcium and triggering the unfolded protein response, leading to apoptosis.

Necrosis can follow more severe elevations of FC. In this situation, it may be cholesterol crystals that induce damage to

showed that SR-A is the main scavenger receptor for group B intracellular organelles to cause necrosis. One response of cells to mitigate the potential toxicity of FC is stimulation of phosphatidylcholine (PC) synthesis and the formation of cellular whorls consisting of FC and PC. If the molar ratio of FC: PC remains above 0.4, necrosis ensues. 266 Limitation of PC synthesis therefore aggravates FC toxicity. In addition, incubation of cholesterol-loaded foam cells with HDL helped prevent necrosis. Deficiency in macrophage-specific ACAT1 in LDLR-deficient mice The history of scavenger receptor knockouts has been led to increased free cholesterol and plaque size, 267 possibly due to controversial. Thus, a recent knockout study of both SR-A (coded an excessive increase in FC: PC. 266 In contrast, overexpression of neutral CEH resulted in increased cholesterol efflux from8 macrophages and decreased atherosclerosis. ²⁶⁸ A key difference appeared to be the lack of upregulated efflux of free cholesterol in 2 the case of ACAT1 deficiency, resulting in toxic accumulation of o free cholesterol. These observations suggest different intracellular trafficking of FC and release after inhibition of ACAT1 compared^{T»} with CEH.

The Niemann-Pick C1 (NPC1) protein appears to act as a shuttle o for FC between cell organelles. Apo E-deficient mice with NPC1 knockout have worse atherosclerosis and, interestingly, severe, 5 spontaneous thrombosis. The cause of the prothrombotic state was 0. not clear. 269

Other genetic models with impaired cholesterol unloading of foam cells show accelerated atherosclerosis. One such model is clearly apo AI deficiency (with protection shown by o overexpression of apo AI). ^{270,271} Severe, fulminant atherosclerosis _f and coronary lesions with appearance similar to human lesions develop in apo E-deficient mice lacking the SR-B1 receptor. 272 Macrophage-specific knockout of SR-B1 had a much lesser effect on atherosclerosis, suggesting overall reverse cholesterol transport (or uptake of HDL-cholesterol into the liver through SR-B1) as perhaps more important than macrophage-specific transport through SR-B1. ²⁷³ Other studies also question the importance of macrophage SR-B1 in reverse cholesterol transport, whereas ABCA1 and ABCG1 do facilitate such transport from cholesterol-filled macrophages (Table 8-4). 274

ABCA1 (the gene found deficient in Tangier disease) facilitates free cholesterol efflux to lipid-poor apo AI discs; the more recently identified ABCG1 can promote efflux to larger, spherical HDL 2 and HDL 3 particles. In addition, ABCG1 promotes efflux of oxysterols. Both receptors appear to be important for macrophage cholesterol efflux and reverse cholesterol transport. 270,271 Strangely, wholebody knockout of ABCA1 in either apo E- or LDLR-deficient mice did not affect atherosclerosis, but serum cholesterol levels were much lower in these mice. On the other hand, macrophage-specific knockout of ABCA1 in apo E-deficient mice did not affect lipoprotein levels but did result in a modest increase in atherosclerosis. ²⁷⁵ In contrast, either whole-body or macrophagespecific knock out of ABCG1 did not consistently increase atherosclerosis (some studies even showed a decrease), possibly because of upregulation through LXR of both ABCA1 and apo Emediated adequate efflux. LXR may also have been stimulated by increased exposure to oxysterols in these ABCG1 knockout mice. However, macrophage-specific knockout of both ABCG1 and ABCA1 resulted in much more atherosclerosis than either ABCA1 or AGCG1 deficiency alone did. ²⁷⁶ These results strongly suggest that both ABCA1 and ABCG1 are important for cholesterol efflux from macrophages in vivo but again demonstrated redundant

Studies generally support the concept that reverse cholesterol transport is atheroprotective. An assay to document

TABLE 8-4	enes Related	to Promotion of Atherosclerosis *
genes	Effect	Gene Function
аро В	FF	Proteoglycan binding-defective apo B tested
LPL (BMT) LRP1	F T	Macrophage LPL can bind lipoproteins
		LRP1 internalization defective knock-in; triglyceride-rich receptor residue
ASM	FF	Acid sphingomyelinase cleaves sphingomyelin to ceramide
sPLA 2, group III	tut	Produces lysophosphatidyl choline in LDL
sPLA ₂ ,group V	F	Produces lysophosphatidyl choline in LDL; BMT study
EP4	FF	Macrophage PGE 2 receptor
5-LO (BMT)	F	Global heterozygotes had much lower levels of atherosclerosis
iNOS (BMT)	F	Produced both excess NO and superoxide in lesions
MSR	NS to FF	Macrophage scavenger receptor; variable subsequent studies
CD36	NS to F	A class B scavenger receptor
Grb2 (BMT)	F	Adapter protein coupling receptor signals to the MAPK pathway
NF- KB , p50 (BMT) F		More macrophages in lesions but absence of foam cells
JNK1	NS	Strongly proinflammatory MAPK
JNK2	FF	Important in macrophages; less foam cell formation in knockout
LXR aP (BMT)	tut	Multiple effects on macrophage cholesterol homeostasis
PPARγ (BMT)	Т	PPAR y /RXR dimer upregulates LXR as well as CD36 transcription
aP2 (BMT)	F	Fatty acid-binding protein found in macrophages and adipocytes
ADFP (BMT)	F	Promotes lipid storage
ACAT1 (BMT)	tut	Catalyzes cholesterol esterification in macrophages
CZECH REPUBLIC (TG)	FF	Transgenic macrophage-specific cholesteryl ester hydrolase
NPC1	tut	Intracellular FC trafficking
SR-B1	tut	An HDL receptor; whole-body knockout showed severe disease
ABCA1 (BMT)	Т	Cellular cholesterol exporter
ABCG1 (BMT)	NS to TT	Increased atherosclerosis when combined with ABCA1 knockout

8

macrophages loaded with radiolabeled cholesterol are injected intraperitoneally into mice. Appearance of the labeled examples of counterintuitive findings are consistent with the and may prove to be the single most frequent general concept of reverse cholesterol transport as measured by this in vivo assay. Perhaps most clearly illustrated is the

seemingly paradoxical finding of increased HDL seen with whole-body or hepatic SR-B1 deficiency. Reverse cholesterol transport through HDL to the liver is clearly impaired in this model. Conversely, overexpression of hepatic SR-B1 lowers HDL but increases reverse cholesterol transport as well as inhibits atherosclerosis. 277

Surprising findings regarding lecithin-cholesterol acyl transferase (LCAT) knockout or overexpression models may also be reconciled by the same technique. Thus, LCAT overexpression may actually increase atherosclerosis and seems to impair wholebody reverse cholesterol transport by decreasing availability of very small, lipid-poor HDL, which are the most active cholesterol acceptors. Conversely, LCAT knockout has been reported to decrease atherosclerosis in some mouse models. Furthermore, in LCAT-overexpressing mice, transgenic expression of cholesteryl ester transport protein appears to facilitate reverse cholesterol transport and to decrease atherosclerosis. 271

PROGRESSION OF ATHEROSCLEROSIS AND PRECIPITATION OF EVENTS

As noted in the introduction, this phase of atherogenesis has to do with the growth of complicated plaques, thrombosis, plaque instability, and precipitation of acute events. Animal models of these more advanced stages of atherosclerosis are relatively few and may be less comparable to the human condition. Indeed, only a few genetic models of atherosclerosis clearly progress to atherothrombotic complications such as myocardial infarction. For example, one model required placing apo E-deficient mice on a high-fat diet after placement of a cast around the carotid artery, giving intraperitoneal injections of lipopolysaccharide to cause immune activation, and producing a hyperdynamic circulation and stress with noise and foot shocks. More than 60% of such animals had carotid plaque ruptures, but thrombosis was still rare because of the vigorous fibrinolytic activity of mice. ^{278,279} In another model, stable transgenic overexpression of human PAI-1 resulted in frequent coronary artery thrombi and myocardial infarction in normolipidemic mice, but this occurred without atherosclerosis. 280 The relevance of this model to the human situation is clearly questionable.

Despite difficulties, advances in this area continue to be made rapidly through direct study of human plaque and pathophysiological studies. Of major importance was recognition of the vulnerable plaque, whose thin fibrous cap, decreased collagen content, and greater inflammatory activity including MMP expression lead to rupture or fissuring in the fibrous cap, exposure of the highly thrombogenic plaque substance to blood, and precipitation of acute thrombotic events. 281-283 Other prominent features of vulnerable plaques include a high free cholesterol content, markers of red blood cell infiltration, and increased vasa vasorum. 283-285

Recently, hypoxia within advanced human atherosclerotic plagues, with release of hypoxia-inducible factors and plague penetration with vasa vasorum, has been related to enlarge ment of lesions. 286,287 Red cells appear to be major contributors of free cholesterol in advanced plaques as well as a source of strongly proinflammatory and pro-oxidant material. Degree of instability has been correlated with degree of neo vascularization of plaques. Whereas the molecular biology of angiogenesis may prove to be

reverse cholesterol transport in vivo was developed in which critical to the progression of advanced human lesions, this topic is beyond the scope of this chapter.

The new observation that abrupt crystallization of choles terol cholesterol in the blood and feces is tracked. 271,277 Several probably contributes majorly to acute events is highly compelling

^{*}Note: many of these studies were performed as bone marrow transplantation (BMT) of the knockout to show that it affected macrophage function specifically NS, not significant; TG, transgenic with overexpression.

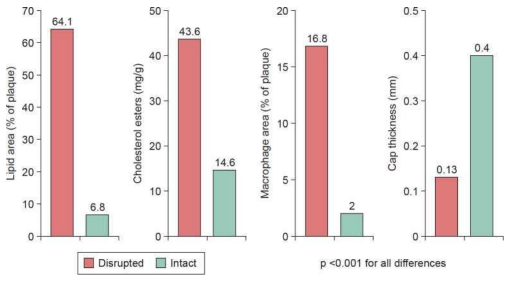


FIGURE 8-10 Composition and morphology versus stability in 668 disrupted or intact aortic plaques from 30 men with CAD. Lipid concentration correlated positively with macrophage accumulation and negatively with minimum cap thickness. (Modified from Felton CV. Crook D. Davies MJ. Oliver MF: Relation of plague lipid composition and morphology to the stability of human aortic plaques. Arterioscler Thromb Vasc Biol 17:1337, 1997.)

cause of event precipitation. ¹⁷ Prior tissue preparation techniques nonsignificantly increased lesion area and promoted increased using ethanol dissolved the crystals and thus pre vented their apoptosis, plaque necrosis, decreased collagen deposition, and proper viewing. Thus, when air dried, almost all lesions associated thinning of the fibrous cap in apo E knockout animals. 297 These with acute coronary events were seen to have penetrating findings are reminiscent of findings, noted previously, of more about by cholesterol crystallization, which may provide the induced by SR-A and STAT1 in already stressed macrophages. underlying force to precipitate fissures and ruptures of the fibrous cap as well. 288,289

With regard to vulnerable plaques, one major paradigm should be kept in mind. Among the strongest correlates of instability (if not the strongest predictor) is lipid content. This was shown strikingly in autopsy studies by Michael Davies' group (Fig. 8-10). ²⁹⁰ All measures of plaque vulnerability including macrophage content, leukocytes in response to similar signaling from cytokines. PDGF degree of leukocyte activation, indices of apoptosis, MMP activity, and collagen content are improved rapidly by marked reduction of receptor is an additional important signaling pathway. plasma lipid levels in animal models, whether by diet manipulation 7,10,291,292 or with statins. 293-295 Remarkably, rapid regression of lesions could also be achieved by infusion of cholesterol acceptors such as injected phosphatidylcholine or phospholipid micelles, which apparently mobilized cholesterol from plaques. 14

More recently, extensive egress of foam cells was shown to occur within days of transplanting atherosclerotic vessels from regulate smooth muscle cell migration. ²⁹⁸ Smooth muscle cellgenetically hyperlipidemic mice into wild-type mice. 14,296 This egress was dependent on signaling in foam cells through the CCR7 chemokine receptor and could be blocked by antibodies against the natural ligands to CCR7, namely, CCL19 and CCL21. Furthermore, expression of CCR7 was increased by LXR agonists. The hyperlipidemic environment, in some way, led to retention of foam cells in the plaque with progression through apoptosis and necrosis followed by formation of the necrotic core. If macrophages maintain their ability to egress freely from lesions, their entry and egress may enhance plaque regression. 14 Interestingly in this considered to stabilize human lesions. They produce collagen in regard, activation of the MAPK p38 can promote cell survival in response to PDGF and TGF-\$, which would be expected to protect macrophages (much as ERK1/2 does in endothelial cells through activation of Akt); activation of JNK promotes apoptosis, especially when endoplasmic reticulum stress is present. This prosurvival effect of p38 may depend on free cholesterol content and other signals (resulting in a net apoptotic signaling from p38 in some knockout of macrophage Nevertheless,

Molecular Biology and Genetics of Atheroscleros cholesterol crystals. There is a distinct volume expansion brought advanced later complications with greater macrophage apoptosis 🖁

Smooth muscle cells are important components of plaques, particularly advanced plaques. Whereas smooth muscle cells can be converted to foam cells, their presence in the plaque is generally considered to promote stability, as they form the fibrous cap and are the source of most collagen and elastin in the plaque. In general, they migrate from the media into the intima like monocytes or other (which exists in iso forms AA, BB, and AB) acting on the PDGF

Both plasmin and MMP-9 are important in facilitating smooth muscle cell movement through the extracellular matrix. Urokinasetype plasminogen activator (u-PA) activates plasmin release by smooth muscle cells, which activates MMPs, particularly MMP-9. Interestingly, both the PDGF receptor and a u-PA receptor interact with surface lipoprotein receptors (LRP-1, LRP-1B, LR11) that specific knockout of LRP-1 greatly increased atherosclerosis in LDLR knockout mice by removing a strong suppressive effect of this receptor on the PDGF receptor (which action is independent of any lipoprotein binding). 299 In contrast, LR11 appears to facilitate u-PA signaling and smooth muscle cell migration and is downregulated by statins. ²⁹⁸ These findings suggest that at least to some degree, smooth muscle cells can contribute to the growth of plaques.

Despite these findings, smooth muscle cells are generally against rupture. TGF- is anti-inflammatory as well as profibrotic. Interruption of TGF- \$\signaling\$ signaling resulted in unstable plaques in apo E-deficient mice. 300,301 Suppression of smooth muscle cell production of collagen by IFN -γ from T cells is generally considered destabilizing. 12 Localized



108 overexpression of p53 in carotid plagues of apo E-deficient mice led to an increase in smooth muscle cell apoptosis in the fibrous caps, thinning of the cap, and spontaneous rupture on hemodynamic challenge with phenylephrine. 302 In addition, activated macrophages, which are associated with unstable plaques, can produce FasL, which binds to Fas (the so-called death receptor) on smooth muscle cells, promoting smooth muscle cell apoptosis. 303 This may help explain plaque instability associated with increased macrophage numbers.

8 If the role of smooth muscle cells in advanced plaques seems somewhat puzzling, the potential impact of proteolytic enzymes on extracellular matrix and atherosclerosis is even less predictable. All migrating cells secrete various proteolytic enzymes to enable movement through the extracellular matrix. Such migration includes not only the translocation of inflammatory cells into the intima but their egress as well. Thus, movement of many cells, even smooth muscle cells, in or out of the plaque might leave it weakened.

Virtually all MMPs and cathepsins are expressed in advanced human plaques. 12 Furthermore, extracellular matrix remodeling, compensatory arterial enlargement, plaque angiogenesis, and plaque disruption will all be affected by proteolytic MMPs, tissue inhibitor of MMP (TIMP), and the highly active cathepsins. Formation of penetrating vasa vasorum seems to be stimulated by MMP-9. 304 Some effects of these various enzymes on atherosclerosis are briefly reviewed in Table 8-5. However, the interpretation of findings in animal models of advanced atherosclerosis remains controversial.

An additional feature of smooth muscle cell behavior in advanced plaques is surprising. Intravascular ultrasound investigation of culprit plaques causing acute coronary syndromes in humans shows that plaque rupture typically occurs just proximal to (upstream of) the point of maximum stenosis. This is the area of most rapid flow. 306 This is true even though the area of slow flow just downstream of the stenosis is where the endothelium is activated, where monocytes penetrate, and where foam cells accumulate with resulting growth of the plaque. It appears that whereas high shear stress promotes endothelial integrity and higher NO production in this upstream area, the higher shear may promote plasmin production as well. Furthermore, underlying smooth muscle cell proliferation and collagen synthesis may be suppressed by the increased NO release. In addition, exposure of activated macrophages to NO may actually stimulate MMP release and promote smooth muscle cell apoptosis. Such changes may be involved in normal compensatory vessel enlargement in areas of rapid flow, particularly during embryogenesis; but in the context of an advanced atherosclerotic plaque, they lead to paradoxical weakening of the fibrous plaque just under the area of endothelium exposed to the highest flow rates. 307,308

Clotting and Atherosclerosis

hypotheses about genes related to thrombosis to be properly thought. 313 tested, and there are many technical difficulties in the for the two conditions are also different.

In human studies, genetic factors clearly increasing the risk result in any bleeding of venous thromboembolism do not necessarily increase the risk for

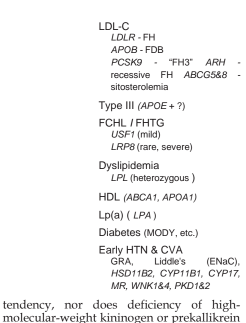
TABLE 8-5		ed to Progression of Advanced Atherosclerosis and Precipitation of Acute Events
genes	Effect	Gene Function
IKK2 (BMT)	Т	IKK2 activates NF - KB activation
p38 (Cre-lox) · LRP-1	NS tut	A MAPK; antiapoptotic in macrophages
		Cre-lox knockout in smooth muscle cells only
MMP-1 (TG)	4 32%	Macrophage-specific TG; mice do not have MMP-1
MMP-3	tut	Also known as stromelysin-1
MMP-7	NS	A matrix metalloproteinase
MMP-9	tut	Also known as gelatinase B
MMP-12	44	MMP-12 may be critical for macrophage migration
TIME-1	NS	Tissue inhibitor of metalloproteinases
CatK	4	Cathepsin K, an elastase
CatS	4	One of the most potent elastases known
Cystatin C	NS	An endogenous inhibitor of cathepsins
IP	Т	PGI ₂ receiver; vasodilating, inhibits platelet activation
TP	44	TXA ₂ receptor; vasoconstrictor, activates platelets
VWF	4	Facilitates both platelet and macrophage adhesion
TFPI	Т	Blocks active tissue factor
V-Factor	Т	FV Leiden mutation tested
Factor VIII	4	A coagulation factor
TM	Т	Thrombomodulin activates protein C
Fibrinogen	NS to 444	Major effect only in human Lp(a) TG mice
Plasminogen	tut	Precursor to plasmin
HCII	Т	Heparin cofactor II is an endogenous thrombin inhibitor
PAI-1	NS	A tissue plasminogen activator inhibitor

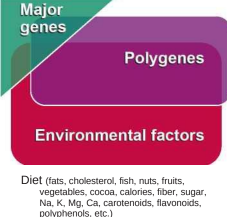
The Cre-lox protocol in this study allowed knockout in macrophages only.

BMT, bone marrow transplantation; NS, not significant; TG, transgenic with overexpression

arterial thrombosis, with only slightly increased but significant odds ratios for myocardial infarction associated with factor V Leiden and Whereas thrombosis clearly plays a role in the progression of prothrombin 20210A in a very large meta-analysis. 311 Mothers with advanced, complex plaques, resulting in demonstrable rapid sons having hemophilia (essentially all heterozygotes) had progression of obstructive lesions, 309,310 genetic predisposition measurably increased risk for bleeding episodes but only an to venous thrombosis does not necessarily coincide with a estimated 36% decreased risk of ischemic heart disease death. 312 predilection to atherosclerosis. Furthermore, only a few Nevertheless, deficiency of von Willebrand factor (vWF) and other animal models develop sufficiently advanced disease for forms of hemophilia appear to be less protective than was once

A full consideration of the molecular biology of thrombosis is interpretation of these models. ²⁷⁹ In arterial thrombosis, clots beyond the scope of this chapter. However, the classic models of soare rich in platelets and poor in fibrin; in venous thrombi, - called intrinsic and extrinsic pathways were based on in vitro assays platelets are broken, fibrin predominates, and the clots are that have little to do with in vivo hemostasis. Thus, complete friable and prone to large emboli. Well-recognized risk factors deficiency of factor XII (Hageman factor, the initiating factor of the so-called intrinsic pathway through contact with glass) does not





polyphenols, etc.)

Oral contraceptives Cigarettes Stress, anger Inactivity Sleep Air pollution

ABCA1, APOA1, LPL, LIPC, LIPG, CETP, GALNT2, ANGPTL4, TRIB1, MMAB BP (AGT, adducin, etc.) Type 2 Diabetes TCF7L2, IGF2BP2, FTO, 9p21.3, KCNJ11, PPARG, CDKAL1, IDE, SLC30A8 BMI (FTO) **Thrombosis** KIF6? FV, prothrombin 9p21.3 markers TNFSF4?

homozygous expression of factor V Leiden or knockout of thrombomodulin, both models of impaired protein C activity and hence increased thrombin effect, led to somewhat larger but much more stable lesions in apo E knockout mice. 320 Other seemingly conflicting results regarding seemingly conflicting results regarding thrombosis have been reported. Thus, plasminogen deficiency markedly increased atherosclerosis in apo E-deficient mice 321 but decreased atherosclerosis substantially in a model of u-PA

FIGURE 8-11 A summary of human genetic and environmental risk factors for atherosclerosis. Major genes linked to atherosclerosis are listed on the left; polygenes (genes have small, presumably cumulative effects) are listed on the right. Environmental factors are listed in the center

XII have a tendency towards thrombosis, possibly through a role in activate u-PA and thereby to promote macrophage activation. 322 activating fibrinoly sis. 314 Also, these cascades fail to depict the critical participation of platelets and tissue factor-bearing cells in thrombosis. Tissue factor is expressed not just on activated PARALLELS WITH HUMAN GENETICS macrophages but also on synthetic smooth muscle cells in atherosclerotic plague. 315 Rather than having two possible cascades, all clot ting in vivo proceeds essentially by a single, highly regulated pathway through exposure to tissue factor-bearing cells. 316,317

Whereas activation of thrombin and subsequent thrombo sis in the face of plaque rupture can obviously be expected to have adverse consequences in acute coronary syndrome, the effect of the various elements of the clotting cascade on atherogenesis at earlier or even to be clinically useful. 326-328 Other variants were found less and less later stages is not straightforward or predictable (see Table 8-5). predictive as better data became available. 329-337 Many of these Thrombin, although clearly the central player in thrombogenesis, also activates PAR1, 2, and 4 (protease-activated receptors 1, 2, and 4) and can thereby have various additional effects. For platelets, thrombin signaling through PAR1 and 4 is activating. In contrast, thrombin can induce PAR1-mediated forearm dilation in humans through stimulated release of NO, PGI 2, and endothelium-derived hyperpolarizing factor; yet in the context of more prolonged exposure, at least in vitro, throm bin suppresses eNOS and promotes endothelin release, stimulates fusion of Weibel-Palade bodies with the plasma membrane, promotes expression of adhesion molecules, and generally activates endothelial cells. Thrombin has multiple additional proinflammatory effects and stimulates activation of NADPH oxidase and ROS production. Finally, active thrombin may genome-wide association studies. Major genes, polygenes, directly promote smooth muscle cell migration and proliferation directly through PAR1 on smooth muscle cells or by increasing PDGF release from endothelial cells. 318

Interventions on thrombin have yielded contradictory findings in experimental settings. A direct thrombin inhibitor, melagatran, decreased lesion size and improved lesion stability with thicker fibrous caps, smaller necrotic cores, and decreased MMP-9 in apo Edeficient mice. 319 However,

LDL-C APOB, APOE, HMGCAR, LDLR, PCSK9, SREB-2, SORT1, TRIB1, NCAN Triglycerides ĂPOA5, APOB, LPL, MLXIPL, GCKR, TRIB1, GALNT2, NCAN ANGPTL3, ANGPTL4 HDL-C

(cofactors with factor XII). If anything, persons deficient in factor overexpression by macrophages. The plasminogen was thought to

Our current understanding of the genetics of human atherosclerosis bears little resemblance to the vast and insightful literature reviewed here. Very few CAD associations with candidate gene variants have been consistently replicated. 323-325 Some candidate gene variants seem to persist with very modest associated odds ratios for CAD in large meta-analyses, but probably none can currently be considered disappointing false starts were likely due to false-positive early findings or "winner's curse." However, some of the apparent lack of measurable effects on atherosclerosis may be due, in part, to the redundancy of so many pathways affecting atherogenesis.

Effects of most of the genes that have consistently been associated with human atherosclerosis risk or risk factors are summarized in Figure 8-11. In general, recognizable major genes affect relatively few individuals but have large effects on atherosclerosis risk. Most of the recognized major genes listed in Figure 8-11 affect lipoprotein metabolism. Polygenes are probably common, but their effects have been more difficult to define or to confirm. Most of those listed on the right side of Figure 8-11 have been reported in relatively recent and environmental factors all interact with each other to ultimately CONCLUSION determine risk. On the basis of international comparisons of CAD rates, the environmental factors clearly outweigh genetic factors in Numerous overlapping and redundant pathways are involved in determining overall population risks.

8 clearly warranted at this time. Benefits of such efforts were shown dyslipidemia, atherosclerotic plaques generally do not develop, by our group, which originated the MEDPED program (Make Early even at such predisposed sites. Recognition of multiple lipoprotein Diagnoses to Prevent Early Death), 338 but the real potential for this modifications, including nonoxidative, that can result in both program has been exemplified by the outstanding results of the endothelial activation and foam cell formation may illustrate the Dutch MEDPED program. 339

In contrast to the relatively poor showing for candidate genes, narrowly focused on just one aspect of atherogenesis. large genome-wide association studies with multiple, built-in consistently associated with CAD risk. However, all these may be due, in part, to the redundancy of the pathways involved. this locus). The odds ratio for CAD is approximately 1.25 for atherosclerotic disease. heterozygote carriers and 1.5 for homozygotes. 340 Several other less REFERENCES well replicated loci from genome-wide association studies include 1p13, MIA3, CXCL12, 21q22, PHACTR1, WDR12, 341 SH2B3, 342 and MRAS. 343 Still other variants are supported by associations in several populations and have not yet been clearly refuted, although lack of replication in newer genome-wide association studies is concerning. These include KIF6 (the W719R variant) 344, 345 and possibly TNFSF4. 323, 330, 332

Whereas much has been said about the potential utility of genetic risk scores, a careful family history together with standard risk factors (together with serum lipoprotein(a) and C-reactive protein) currently remains considerably more predictive than available genetic scores. 341,346-349 The potential utility of genetic tests to guide personalized medicine in preventing or treating atherosclerosis has similarly yet to be seen. There is much hope that these limitations may be overcome in the near future. On the other hand, it may be that the strongest associations have already been found (at least for common variants). Indeed, the vast complexity of potential genetic and environmental factors and their interactions, the potential myriad contributions of rare variants, and these all working in the context of extensive redundancy of atherosclerosis-related pathways may preclude our ever being able to make greatly 11. improved prediction of individual disease. Certainly, we must recognize currently our extremely limited ability to predict the 12. presence of atherosclerotic disease in an individual patient before its clinical manifestation.

screening for atherosclerosis may obviate the need for individual risk prediction by risk factors other than for general classification into groups needing earlier or later screening. One can envision a 16. paradigm in which vigorous medical interventions (particularly lipid-lowering medications) would be directed empirically to those with disease found by noninvasive screening and generally 18. withheld from those with no evidence of disease. If this proves to be the most effective management model, insights gained from genetic 19. studies would be more important as a means to clarify pathophysiology in general, to stimulate the development of innovative new therapies, and potentially even to help guide intervention. Such an outcome would arguably still justify our 21. efforts to identify genetic associations in human populations, even if they did not ultimately prove useful for determining individual

initiation, promotion, and progression of atherosclerotic plaques After review of the literature and the additional insights and precipitation of clinical events. Much recent progress has been presented in Figure 8-11, it is sobering to realize that familial made regarding how endothelial cells transduce changes in blood hypercholesterolemia provides the only example of a genetic cause flow into intracellular signals that control the location of sites of premature CAD for which a systematic, population based predisposed to plaque development. However, without the effect of approach aimed at proband identification and family screening is standard risk factors, particularly at least some permissive level of futility of using an intervention (eg, antioxidants) that is too

The difficulty in identifying new genes associated with human replications have begun to provide genetic variants that are atherosclerosis (particularly loci unrelated to standard risk factors) associations are very modest. The most consistently identified of Nevertheless, greater understanding of atherogenesis and these are the variants at 9p21.3 (such as rs1075724 with strong continued progress in the treatment of risk factors will undoubtedly linkage disequilibrium between all the risk-associated markers at lead to improved therapies that will further reduce clinical

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ABBREVIATIONS Used in Chapter 8

Akt). In general, a followed by lowe refer to the huma but the literature common usage is	reviations are traditionally not fully capitalized (abbreviations that begin with an initial capital let r case letters refer to the mouse gene, whereas all r is not consistent in this use. The most generally given here. An r may be added to shown here to indicate the plural (as in	ter	abbreviated from Japanese, <i>ju-nana</i> , the number 17; derived from the ASV 17 provirus; also referred to as simply Juncyclooxygenase 2 cAMP response element cAMP response element-binding protein Creactive protein colony-stimulating factor 1 receptor; also known as FMS; receptor for MCSF cytidine triphosphate:phosphocholine cytidylyltransferase connexin 43
ACAT1	acyl coenzyme A:cholesterol acyltransferase (ACAT) 1; ACAT1 is in macrophages; ACAT2 is in liver and intestines, not macrophages	CT Cx43 DAG	diacylglycerol Dia-interacting protein 1 dedicator of cytokinesis 2 dedicator of cytokinesis 180;
ADAMTS1 ADFP ADMA AGE Act	a disintegrin and metalloproteinase with thrombospondin motif 1 adipose differentiation-related protein asymmetric dimethylarginine advanced glycation end products also known as PKB; first identified as an oncogene from the AKT-8 thymoma cell line, which was derived from the AKR/J mouse	DIP1 Dock2 Dock180 ECM ECSIT	also known as Dock1; abbreviation can also mean "down stream of Crk-binding protein" extracellular matrix evolutionarily conserved signaling intermediate in toll pathways endothelium-derived hyperpolarizing factor epidermal growth factor receptor early growth response 1; a transcription factor extincted by DNA
AP-1	activator protein 1, which contains c-fos and c-Jun antioxidant response element apoptosis	EDHF	activated by JNK engulfment and cell motility gene 1 endothelial nitric oxide synthase
ARE ASK1 ASM ATF2	signal-regulating kinase 1 acid sphingomyelinase activating transcription factor 2; the protein forms a homodimer or heterodimer with c-Jun and stimulates CRE-dependent transcription adenosine triphosphate angiotensin II type 1 receptor Bcl-2-associated death promoter; a key regulatory protein for apoptosis; when phosphorylated (through growth factor receptors), it prevents apoptosis; when	EGFR Egr-1 ELMO1 eNOS EP2 ERK ET-1 FAK Fas	prostaglandin E 2 receptor extracellular signal-regulated kinase endothelin 1 focal adhesion kinase apoptosis-stimulating fragment; Fas is actually a cell receptor Fas ligand fibroblast growth factor receptor 2 filamin A-associated Rho GAP friend murine leukemia virus integration site 2; also known as CSF1R fibroblast growth factor receptor substrate 2 a Grb2-associated binder 1 glycosaminoglycan
R Bad	unphosphorylated, it promotes apoptosis Bcl- 2-associated X protein B-cell CLL/lymphoma 2 bone morphogenic protein 2 or 4 calcium- and DAG-regulated GEF1 calcium/calmodulin-dependent protein	FasL FGFR2 FilGAP Fms	GTPase-activating protein (generally - inactivates Ras family GTPases by converting GTP to GDP) glyceraldehyde 3-phosphate dehydroge nase G nucleotide dissociation inhibitor (inhibits
Bass Bcl-2 BMP2/4 CalDAG-GEF1 CaMKII	kinase II CREB-binding protein (also called p300 homologue or p300 or p300/CBP) chemokine, CC motif, ligand 2 (also known as MCP-1) chemokine, CC motif, receptor 2 cluster of	FRS2 Gab1 GAG GAP	its dissociation of GDP or GTP from Ras-like proteins and maintains them as soluble cytosolic proteins, thereby blocking Ras or Rho signaling) guanosine exchange factor; GEF facilitates the exchange of GTP for GDP to activate Ras
CBP	differentiation 14 CD40 ligand; CD refers to "cluster of		family proteins
CC12	differentiation" cell division cycle 42 (a Rho family GTPase)	GAPDH	G protein-coupled receptor growth factor receptor-bound protein 2 (acts as an adapter
CCR2 CD14 CD40L	neutral cholesteryl ester hydrolase cholesteryl ester transport protein FBJ (Finkel-Biskis-Jinkins) oncogene osteosarcoma; also known as FOS or Fos. The "c" refers to cellular as	GDI	protein between receptors and Sos)
Cdc42	opposed to "v" for viral		
CZECH CETP c-fos		GEF	
		GPCR Grb2	

118 GRO	growth-regulated oncogene a; also known MI		macrophage inflammatory protein 1a; also
	CXCL1GSSG glutathione-glutathione disulfide (oxidized glutathione) glutathione- S -transfera	ase MKK	known as CCL3 macrophage inflammatory protein 1\$; also
CCT	guanosine triphosphate	MKP-1 MLCK	known as CCL4
GST GTP	hydrogen peroxide heparin cofactor II	mmLDL	MAP kinase kinase MAPK phosphatase 1 myosin light-chain
H_2O_2	high-density lipoprotein(s)	MMP	kinase minimally modified LDL matrix
HCII	hydroxyeicosatetraenoic acid	MSR	metalloproteinase macrophage scavenger
HDL	high-mobility group box 1 4-hydroxy-2-nonen		receptor (name of the gene that gives rise to
HETE O HIMORA	oxygenase 1 hypochlorous acid heat shock transcription factor 1 heat shock pro	MyD88	both SR-AI and SR-AII through alternate
8 HMGB1 4-HNE	aids in folding NADPH of nascent proteins; H		splicing) myeloid differentiation primary response
HO-1	promotes eNOS activity; HSP-60 may be expre		protein 88
HOC1	in endothelial stress or activation and promote		nicotine adenine dinucleotide phosphate (H
HSF1	autoimmunity	NF-B NK cells	= reduced form)
HSP	interferon- γ inhibitor of NF- κ B or inhibitor- K B	Nrf2	NF-KB essential modulator (part of the IKK
	IKB kinase interleukin	NO	complex); also known as IKKY nuclear factor-KB
	inositol 1,4,5-triphosphate (soluble cyto-	O 2	natural killer cells
	solic intracellular messenger released after	Oct1	nuclear factor erythroid 2-like related factor
IFN	action of PLC)	OH	2
IB IKK	interleukin-1 receptor-associated kinase 1 interferon regulatory factor 3	OLR1 ONOO -	nitric oxide
IL IL	Janus kinase (part of JAK-STAT pathway)	oxLDL	superoxide anion octomer-binding transcription factor 1 hydroxyl radical
IP3	c-Jun N-terminal kinase (a MAPK) Kelch-like		oxidized LDL receptor 1 gene peroxynitrite
	derived cap'n'collar	PAF	radical
ID 4 1/4	homology-associated protein 1	PAF-AH	oxidized LDL (generally considered rather
IRAK1 IRF3	Kruppel-like factor 2 leukemia-associated Rho known	PAK	severely oxidized) platelet-activating factor
JAK	as ARHGEF12	PAMP	platelet-activating factor acyl hydrolase, also called LpPLA2p21-activated kinase
JNK	low-density lipoprotein(s)	PAR1	pathogen-associated molecular pattern(s)
Keap1	LDL receptor lipoic acid synthase	Par6	protease-activated receptor 1 (the receptor
KLF2	lin-11 isl-1 match-3; the terms <i>lin</i> (for cell linea		activated by thrombin) partitioning defective
LARG	mec have genes in <i>Cae</i> norhabditis elegans that have homologous	-PARG PARP-1	protein 6 (involved in cell polarization with
	PCAF domains	111111 1	Cdc42) poly(ADP-ribose) glycohydrolase poly(ADP-ribose) polymerase 1 p300/CBP-
LDL LDLR	LIM domain kinase	PDGF	associated factor platelet-derived growth
LIAS			factor
LIM			PDZ domain-containing protein 1; PDZ
			domains frequently involve organization of receptors, ion channels, or signaling mol-
			ecules at the inner cell membrane; PDZ
T T3 47/			stands for the original recognition of shared
LIMK LOX-1	lectin-like oxidized LDL receptor 1; PDZK1 co	dad by	structure in 3 proteins, PSD-95, DlgA, and
LOX-I	the <i>OLR1</i> gene (oxidized LDL receptor 1) lipor		ZO-1
	lipase	rotein	platelet and endothelial cell adhesion mol- ecule 1
LPL	lipoprotein-associated phospholipase A2; same	e as PAF-	prostacyclin pleckstrin homology (a domain
$LpPLA_2$	AH		found in many proteins)
LRP1	LDL receptor-related protein 1		phosphatidylinositol 3-kinase
MAD	mothers against decapentaplegic	PECAM-1	phosphatidylinositol 3,4,5-phosphate (membrane bound)
MAPK	mitogen-activated protein kinase	I LC/IIVI I	Pak-interacting exchange factor beta; also
MAPKK	MAPK kinase, also called a MEK (MAPK/	PGI ₂	known as Cool-1 and Gef7
MAPKKK	ERK kinase)	PH	protein kinase B (also known as Akt) protein
MARCO	MAPKK kinase or MEKK (MEK kinase) macro receptor with collagenous	pnage PI3K	kinase C; there are several different isoforms:
MCP-1 M-	PIP3 structures	11310	a, \$1, \$2, and y are <i>conventional</i> (sensitive to calcium and DAG); S, £, n, and 0 are <i>novel</i>
CSF	monocyte chemoattractant protein 1 monocyte	colony-	(sensitive to DAG but not to calcium); Z, I,
MD 2	stimulating factor; also	-PIX	and X are atypical (not sensitive to calcium or
MD-2	known as CSF-1 origin of abbreviation obscure		DAG) paxillin-kinase linker; also known as
	referred to as lymphocyte antigen (LY) 96; ostr	ici- i KC	GIT2
	tural homology to mite allergen Der p 2 micro	tubule-	
mDia1	organizing formin diaphanous 1		
MEF2C	myocyte enhancer factor 2C MAPK/ERK kinase = MAPKK		
MEK	MEK kinase	PKL	
MEKK	macrophage migration inhibitory factor		
MIF	-		

Src homology collagen-like; an adapter **119** protein phospholipase A₂ (soluble, sPLA₂, and Shc lipoprotein-PLA₂ whose SH2 domain binds phosphorylated tyrosine on associated, LpPLA2, are commonly described, but other the cytosolic domain of RTKs (receptor tyrosine kinases) forms are also known) and also binds Grb2 phospholipase C; different isoforms include \$1-4 SH2 domain-containing tyrosine phosphatase (mainly activated by the Gq SHP-2 **PLC** SMAD proteins are homologues of both the *Drosophila* class of G proteins that couple to GPCRs; protein mothers against decap-entaplegic (MAD) and y1, 2 (activated by receptor protein tyrosine **SMAD** the *C. elegans* protein SMA; the name is a combination of kinases or nonreceptor protein tyrosine kinases and 8 the two smooth muscle cells suppressors of cytokine further stimulated by PIP3); signaling superoxide dismutase (Cu/Zn SOD = \$1-4 (possibly activated by calcium or G copper/zinc superoxide dismutase – in cytoplasm; MnSod = manganese superoxide dismutase—in proteins); £ (activated by Ras and Rap); and Z (found mitochondria) son of sevenless; may be referred to as only in mammalian sperm and **SMC** mSos; a GEF for Ras family proteins; binds Ej to the sensitive to calcium) **SOCS** adapter Grb2 scavenger receptor A (implies both AI and **PLD** phospholipase D SOD AII, alternate splice isoforms of the same MSR gene) **PLTP** phospholipid transfer protein peroxisome proliferatoractivated receptor (there are a, y, and S forms) **PPAR** scavenger receptor B1 (an HDL receptor) © pronounced peroxiredoxin 1 Sos PRDX1 "sarc" (short for sarcoma), the first identified of a family PSGL1 of proto-onco-5-genic tyrosine kinases scavenger P-selectin glycoprotein ligand 1 phosphatase and **PTEN** receptor expressed by endothelial cells scavenger tensin homologue Rac1 receptor-phosphatidyl serine and oxidized lipoprotein Ras-related C3 botulinum toxin sub- SR-A strate 1 signal transducer and activator of transcription 1 spleen murine leukemia viral oncogene homo-RAF1 tyrosine kinase TAK1-binding protein 1 and 2 TGF- \$ -activated kinase logue 1; the derivation of the abbreviation SR-B1 is not 1 (a MEKK) tert -butylhydroquinone TANK-binding clear; a MAP kinase kinase kinase or Src MEK kinase; kinase 1 tissue factor typically activated by Ras but has multiple tissue factor pathway inhibitor transforming growth phosphorylation sites that can upregulate or downregulate factor- \$ triglyceride-rich lipoproteins (includes activity; SREC inactivated when bound to the 14-3-3 reguchylomicrons, VLDL, and remnants) T-helper type 1 **SR-PSOX** latory protein lymphocytes recombinase activator gene 1 receptor for advanced T-lymphoma invasion and metastasis 1 toll-like receptor glycation end prod-STAT1 Ra -4 thrombomodulin tumor necrosis factor- a ucts (AGE) TNF receptor-associated factor 6 TRIF-related adapter Ras-like protein; also RalA Syk molecule tumor necrosis factor receptor-associated regulated on activation normal T cell **TAB1/2** protein toll/IL-1 receptor domain-containing adapter expressed and secreted; also known as TAK1 inducing interferon- \$\bigsep\$ thioredoxin thioredoxin-CCL5 **tBHQ** interacting protein ubiquitin-conjugating enzyme 13 Ras-related protein 1 TBK1 ubiquitin-conjugating enzyme E2 variant 1 isoform A named for a transforming oncogene found TF in rat Rap1 urokinase-type plasminogen activator vasodilatorsarcoma; the Ras superfamily are TFPI Ras monomeric GTPases that act as off-on TGF- switches (off stimulated phosphoprotein (with functions analogous to WASp) named for *vav*, the sixth letter in the Hebrew when GDP is bound, on when TGRL GTP is bound); alphabet (for the sixth oncogene discovered at the members of the family include Ras, Rho/Rac, Rab, Rap, laboratory that characterized the protein); the Vav Arf, Ral, Th1 Ran, Rheb, Rad, Rit, and Miro; more than proteins have GEFs for Cancer TIAM1 a hundred members are known; Ras super-TLR4 family proteins have geranyl-geranyl lipid TM anchors for attachment to the inner cell TNF- membrane Ras homologous; one of the members of **TRAM** the Ras superfamily of GTPases; RhoA, TRAP Cdc42, Rac1 are common Rho family members Rho Rho-associated, coiled-coil containing protein kinase; also referred to frequently as Trx simply Rho kinase TXNIP sphingosine 1-phosphate; receptors are **UbC13** S1P1 through 5 (also known as S1PR1-5 **UEV1A** and "endothelial differentiation gene" or EDG receptors) uPA ROCK stromal cell-derived factor 1; also known VASP S₁P SDF1

as CXCL12

glutathione (the -SH refers to reduced Vav1-3 sulfur on

Src homology domain 2 (a conserved protein domain

that recognizes phosphorylated tyrosines)

SH

SH₂

120 VCAM-1 VE-
cadherin
VEGFR2vascular cell adhesion molecule 1
vascular endothelial cadherin
vascular endothelial growth factor receptor
2, also called Flk-1vWF
WASp
WAVE
WAVE
VAVE
VWASp family verprolin homology domain-
containing proteinVLDLvery-low-density lipoprotein(s)ZO-1zona occludens 1

Blood Pressure

CHAPTER 9

Hypertension: JNC 7 and Beyond

William J. Elliott

139/80-89

- people with chronic kidney disease, or those with established heart disease, but the evidence for these targets is controversial.
- · Despite the widespread availability of many effective and inexpensive antihypertensive drugs, hypertension control rates are

Elevated blood pressure (BP) is responsible for

more deaths than any other risk factor for

cardiovascular disease (CVD), which is

already the leading cause of death and

disability in the developed world and is

expected to become the leading cause of death

and disability worldwide by the year 2025. 1

Historically, "hypertension" (BP > 140/90 mm

Hg) was thought to be the major driver of

these problems, but BP levels between 120-

prehypertension) not only is more prevalent

but also elevates cardiovascular risk. 1,2

Compared with many other risk factors for

stroke, acute myocardial infarction, and heart

failure, hypertension is among the simplest to

diagnose, has the widest variety of treatment

options, and (particularly in high-risk individuals) is the most cost-effective

preventive strategy. 3-5 Because of its high

prevalence (eg, ~ 29% of adults; Fig. 9-1) in the

United States, 6 hypertension ranks first

among the chronic conditions for which

Americans visit a health care provider. 3,7,8 One

of the major reasons for the impressive

reduction in age-adjusted stroke mortality (~

62%) and coronary heart disease (CHD)

mortality (~ 45%) in the United States since

1972 is the wide spread acceptance of the need

to treat hypertension and our increased ability

to reduce BP effectively. 8

Hg

(now

mm

suboptimal. A concerted effort by the health care provider, patient, and health care delivery system is required to control blood pressure in the long term and to reduce the risk of cardiovascular disease.

KEY POINTS

- Reducing high blood pressure, factor, prevents of delays the an important and the most common cardióvascular risk cardiovascular disease
- Measurement of blood pressure as an independent risk factor for cardiovascular disease can be easily and inexpensively accomplished in the medical office, in the home, or by sophisticated devices.
- Dietary sodium restriction reduces the long-term risk of cardiovascular disease, but weight loss is the most effective short-term lifestyle modification to lower blood pressure.
- Drug therapy for hypertension reduces the risk of cardiovascular disease more than placebo or no treatment. Different expert panels do not agree on a universally applicable treatment algorithm, but all agree that a hypertensive drug that improves prognosis can be given to the patient with a compelling indication for that drug.
- The traditional treatment target for blood pressure is <140/90 mm Hg in uncomplicated hypertensives; <130/80 mm Hg has been recommended for diabetics,

HYPERTENSION GUIDELINES

Because of the global public health impact of hypertension, many countries organizations have developed guidelines for its diagnosis and treatment. 3-5,9 There are major differences in how frequently these are updated, and there is little consensus among them regarding risk stratification, initial BP targets therapy, or (Table these Nevertheless interesting recommendations summarize our current knowledge but come to very different conclusions.

PATHOPHYSIOLOGY OF **HYPERTENSION**

Although several genetic forms hypertension have been discovered and many secondary causes of hypertension identified, most hypertensive individuals have primary (or essential) hypertension, for which no specific cause can be found. Many neuro hormonal systems affect BP, and many of have been manipulated pharmacologically to assist with its control. Perhaps chief among these are the renally regulated "BP-vascular volume" concept, the renin-angiotensin-aldosterone (RAAS), and the sympathetic nervous system (SNS). 10 Much recent work has developed the concept of dysregulation of nitric oxide

122 (and oxidative stress) as a potential contributor to CVD, because most hypertensives (especially those older than 55 years) to elevated BP.

DEFINITION AND CLASSIFICATION OF **HYPERTENSION**

From 1974 to 1993, hypertension in the United States was defined only by diastolic BP (> 90 mm Hg) and was classified as mild, moderate, or severe. Since then, burgeoning evidence from both epidemiological studies ¹² and clinical trials ^{13,14} has CVD and renal events than diastolic or pulse pressure is. CVD events, if left untreated. Therefore, the focus has shifted to systolic BP, particularly

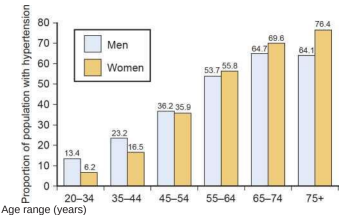


FIGURE 9-1 Age- and gender-specific prevalence of hypertension in the National Health and Nutrition Examination Survey (NHANES) 2005-2006, 8 As in all previous NHANES data sets going back to 1971, hypertension was defined as BP > 140/90 mm Hg or taking antihypertensive medication. The overall prevalence of hypertension was about 29% (no significant change since NHANES 1999-2000), but the prevalence of the awareness, treatment, and control (to < 140/90 mm Hg) of hypertension has improved to 78%, 68%, and 44%, respectively. 6

including hypertension. 11 Although animal studies, - have pretreatment systolic BP > 140 mm Hg rather than diastolic BP epidemiological evidence, small cohort trials, and a few larger > 90 mm Hg. Similarly, the classification system has evolved into ran domized clinical trials have implicated nitric oxide, we "stages" of hypertension (Table 9-2), of which JNC 7 recognizes only have few pharmacologic agents specific to this system that are two: stage 1 (systolic BP 140 159 mm Hg or diastolic BP 90-99 mm clinically useful in large populations of free-living Hg) and stage 2 (systolic BP > 160 mm Hg or diastolic BP > 100 mm hypertensive patients. Further research is therefore necessary Hg). This scheme is independent of gender and age, although many before the Science Magazine "molecule of the year" in 1998 authorities have suggested that higher risk individuals (eg, African achieves the status of the RAAS or SNS as a major contributor Americans, diabetics, kidney and disease patients) should have BPlowering treatment initiated at threshold BPs lower than 140/90 mm Hg. This suggestion has been carried to its logical extreme by several sets of guidelines ^{5,9} that base diagnosis and treatment decisions on the absolute risk of CVD for a given individual and not on any specific cutoff for BP. Proponents of the "polypill" (which contains three antihypertensive agents at moderate doses) have recommended that not only classification of BP but also measurement of BP should be abandoned in favor of BP lowering in demographically defined populations at high risk for CVD. 15 This approach denies the benefits of therapy to those who are likely to demonstrated that systolic BP is a better predictor of future suffer target organ damage and worsened hypertension, rather than

> Prehypertension was defined in JNC 7 as BPs between 120-139/80-89 mm Hg, ³ but the term has been largely ignored by other guidelines. Some believe that the term is too pessi mystical and deterministic, but in the Framingham cohort, more than 90% of individuals who are not already hypertensive at the age of 55 years become so during the next 25 years of follow-up. 16 Individuals with this level of BP clearly have increased CVD risk compared with normotensives (BP < 120/80 mm Hg). ² In the most recent population-based survey of American adults, prehypertension (43% prevalence in men, 30% in women) was more common than hypertension (only 30% and 28%, respectively). ⁶ Because there are many more prehypertensive than hypertensive individuals in most populations, 1 the burden of BP-related illness would soar if this "disease label" was widely accepted. Although lifestyle modifications are generally recommended for these individuals, ³ and pharmacologic therapy with a moderate dose angiotensin receptor blocker (ARB) was "feasible" (and significantly prevented the transition to frank hypertension), ¹⁷ outcomes from large clinical trials are currently lacking to address the question of whether lowering of BP in prehypertensives is effective in preventing CVD. The intent of JNC 7 in introducing the term was to highlight the elevated CVD risk and to motivate affected individuals to pursue strategies to lower their BP, not (as some have suggested) to increase the sales of antihypertensive drugs.

TABLE 9-1	Recommendations of Differ	ent Sets of Hypertension Guideline	s	
	JNC 7, 2003	ASHWG, 2005	NICE, 2006	ESH/ESC, 2007
Risk stratification	Based on BP only	Based on tests for target organ damage, not BP	Based on absolute risk	Extensive system based on absolute risk
Default initial therapy	Low-dose diuretic		Calcium channel blocker or angiotensin- converting enzyme inhibitor (age dependent)	Does not matter; most need two drugs anyway
Beta blockers	Second-line	No comment	Fourth-line	Not with diuretics in patients with metabolic syndrome
BP targets	<130/80 mm Hg for diabetics, chronic kidney disease		Lower for diabetics	<130/80 mm Hg for diabetics, even lower if renal dysfunction and proteinuria >1 g/day

JNC 7, Seventh Report of the Joint National Committee on Prevention, Detection, Evaluation, and Treatment of High Blood Pressure³; ASHWG, American Society of Hypertension Writing Group9; NICE, National Institute for Health and Clinical Excellence (British National Guidelines)4; ESH/ESC, European Society of Hypertension/European Society of Cardiology.5

TABLE 9—2 Some Useful "Cut Points" for Blood Pressure				
		Setting		
JNC 7 Threshold	Office ³	Home (self) 19	24-Hour Ambulatory ²³	
Prehypertension	120/80 mm Hg 115/75	5 mm Hg About 114/		
			72 mmHg	
Target for "high-risk" patients	130/80 mm Hg 130/80) mm Hg Probably 12:	3/ 77 mm Hg	
Stage 1 hypertension (for diagnosis)	140/90 mm Hg 135/85	5 mm Hg 133/82 mm I	Нg	
Stage 2 blood pressure (for diagnosis)	160/100 mm Hg 155/9	95 mm Hg About 152/	93 mm Hg	

Modified from JNC 7, 3 the American Heart Association Scientific Statement, 19 and the European Society of Hypertension. 23

BLOOD PRESSURE MEASUREMENT

favor of its lowering has recently been made, 15 terminology first introduced by Nicolai Korotkoff in 1905 is still used in recording of indirectly determined BPs. ¹⁸ Systolic BP is recognized when clear sounds in the traditional fashion. The oscillometric technique and repetitive tapping sounds are heard; diastolic BP is recorded measures biophysical oscillations of the brachial artery, which are when the sounds disappear. Only when audible sounds are heard down to 0 mm Hg is the "muffling" of sound (Korotkoff phase IV) with a mercury sphygmomanometer; systolic BP is determined recorded, between the systolic reading and zero (eg, 178/72/0 mm directly from the threshold oscillation, mean arterial pressure is Hg).

Techniques of Measuring Blood Pressure

The proper technique of accurate BP measurement is typically taught very early during medical training but rarely followed thereafter. Experience in many clinical trials has shown that retraining and at least annual certification in BP measurement are often required to obtain meaningful BP data. One reason for including a placebo arm in registration trials of new antihypertensive drugs is to account for the many potential confounders in BP measurement, including observer expectation bias. The current push for "pay-for-performance" (which includes BP as one indicator of quality of care) is likely to lead to an overabundance of "8" as the terminal digit preference of BPs recorded by health care professionals in many office settings to ensure the greatest possible proportion of customers who are below thresholds. Interestingly, current Healthcare Effectiveness Data and Information Set (HEDIS) guidelines have made achievement of BP goals more likely. Before 2007, the lowest recorded BP (systolic and diastolic) obtained simultaneously at an office visit was accepted as "the" BP for that visit. In the most recent revision, however, the lowest recorded systolic and the lowest recorded diastolic (even if obtained many minutes apart) are accepted, biasing the probability of a below-threshold BP higher.

Home Blood Pressure Measurements

Because of these and many similar challenges, more emphasis is being placed on measurements outside the medical office. Home (or self) BP monitoring is the least expensive and most widely applicable to large populations. Many convenient, inexpensive, and relatively accurate machines are now available. Some authorities think that such devices should be provided to every person with elevated BP and that their physicians should be paid for interpreting home BP data, ¹⁹ but others are concerned about their widespread use because clinical trials have rarely based their treatment decisions solely on home readings.

Home BP readings are typically lower than measurements taken

in the traditional medical environment (by about 12/7 mm Hg on average), even in normotensive subjects. Home readings are better correlated with both the extent of target organ damage and the risk of future CVD events or mortality than are readings taken in the health care provider's office. Home readings can also be helpful in evaluating symptoms suggestive of hypotension, especially if they are intermittent or infrequent. During treatment, reliable home readings can lower costs by substituting for multiple visits to health care providers.

Current recommendations advocate the use of validated oscillometric devices with an appropriately sized cuff around the upper arm. The device should be calibrated against a standard _ sphygmomanometer (using a Y tube) and the technique of the measurer checked. At least 12 (a week of twice-daily duplicate or⁹ triplicate) readings are the minimum on which to base treatment decisions. 19 Home BP monitoring, coupled with remote monitoring and feedback by a health care professional, has improved BP control rates, perhaps by improving medication adherence. ^{20,21}

Ambulatory Blood Pressure Monitoring

Automatic recorders are now available that measure BP frequently in a 24-hour period, during a person's usual daily activities in a 24-nour period, during a person's usual daily activities (including sleep). In the United States, devices that measure BP Although the recommendation to eschew measurement of BP in indirectly (ie, without arterial cannulation) use either an auscultatory or an oscillometric technique. The auscultatory type 8 uses a microphone placed over the artery to detect Korotkoff a compared (by use of a proprietary algorithm) with those observed estimated, and diastolic BP is calculated. Both types of monitors are light weight (< 450 g), simple to apply and to use, accurate, relatively quiet and tolerable, and powered by two to four small batteries. Data from 80 to 120 measurements of BP and pulse rate (usually every 15 to 20 minutes during waking hours and every 30 minutes during the night) are typically stored in a small microprocessor and then downloaded into a desktop computer, which then edits the readings and prints the report. Much research with the use of ambulatory blood pressure monitoring (ABPM) has suggested that this technique is the most accurate method of measuring BP, correlates most closely with target organ damage, and best predicts future cardiovascular events (even independently of office BP measurements). ²² In the research setting, ABPM readings have therefore become accepted as the "gold standard" of BP measurements. Accordingly, several expert panels have provided both correlations between ABPM results and BP measurements in other settings (see Table 9-2) recommendations for its use (Box 9-1). 22,23 Although this method of measuring BP is becoming "standard of care" in many settings, its use is limited in the United States by restrictions on reimbursement for the procedure, which currently in the Medicare age group amounts to \$60 to \$90, only when the result is a new diagnosis of white coat hypertension.

> ABPM makes it possible to measure BP routinely during sleep and has reawakened interest in the circadian variation of heart rate and BP. Most normotensives and perhaps 80% of hypertensives have at least a 10% drop in BP during sleep compared with the daytime average. Although there may be some important demographic confounders (African



Diagnosis and Prognosis

Evaluation of suspected white coat hypertension

Evaluation of refractory or resistant hypertension Evaluation of circadian pattern of blood pressure

Symptoms

Evaluation of dizziness, presyncope, and syncope

Evaluation of the relationship of blood pressure to clinical symptoms

Evaluation of Antihypertensive Agents (research based)

Evaluation of trough-to-peak ratios (and determine optimal dosing intervals)

Evaluation of antihypertensive efficacy

Evaluation of effects of timing of dosing of antihypertensive agents

Modified from Pickering et al. 22 and O'Brien et al. 23

Americans and the elderly have less prominent "dips"), several prospective studies have shown an increased risk of cardiovascular events (and proteinuria in type 1 diabetics) among those with a nocturnal "nondipping" BP or pulse pattern. Several Japanese studies have raised concern that elderly persons with more than a 20% difference between nighttime and daytime average BPs ("excessive dippers") may suffer unrecognized ischemia in "watershed areas" (of the brain and other organs) during sleep if their BP declines below the autoregulatory threshold.

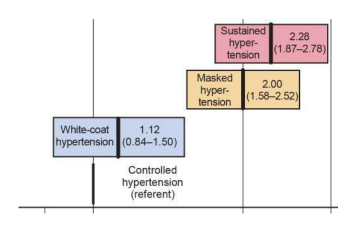
White Coat and Masked Hypertension

Approximately 10% to 20% of American hypertensives have substantially lower BP measurements outside the health care provider's office than in it (so-called white coat hypertension). The white coat itself is unlikely to be the only factor that increases BP. Careful studies originally done in Italy (and now corroborated elsewhere) show that BP rises in response to an approaching physician who is not previously known to the subject. The acute elevation in BP is apparently less marked if a nurse approaches the are significantly lower than those measured in other settings. subject, even if the nurse is wearing a white coat. The Originally thought to be more common in young women with large pathophysiological and psychological "reasons" for this exaggerated BP response are unclear.

The clinical consequences, prognostic significance, and amenability of white coat hypertension to drug treatment are controversial. One point of view suggests that if a person has an acute rise in BP due to "stress" from an approaching physician, similar elevations in BP are likely whenever any stress ful stimulus is encountered. In several convenience samples and populationbased studies, people with white coat hypertension had a greater prevalence of subclinical risk factors for CVD, including left ventricular hypertrophy, family history of hypertension and heart disease, hypertriglyceridemia, ele vated fasting insulin levels, and hypertension as this is the only way to detect masked hypertension. lower HDL-cholesterol levels.

A minority view, based on more conservative definitions of the "white coat effect," proposes that some individuals consistently BLOOD PRESSURE AND CARDIOVASCULAR RISK show a similar and marked elevation in BP in response to the health care environment. Several long-term observational studies have Initial estimates of how much elevated BP increases the risk of heart shown a much reduced risk of either target organ damage or major attack, heart failure, stroke, and other CVD events were derived CVD sequelae among people with lower BPs measured either at home or by 24-hour BP monitoring compared with measurements known of these in the United States is taken in the same person in the physician's office. A recent metaanalysis of the seven published observational studies showed that the 1550 patients with white coat hypertension had a significantly lower risk of CVD than the 4819 with sustained

FIGURE 9-2 Comparison of cardiovascular risk in white coat and masked hypertension with sustained and controlled hypertension. Each box corresponds to the 95% confidence limits (also given numerically within each box), and the dark vertical line within each box corresponds to the adjusted relative risk (also given numerically within each box). This meta-analysis summarizes the 912 first cardiovascular events in seven published observational studies involving 11,502 subjects



with an average baseline age of 63 ± 6.5 years, during 8.0 years of mean follow-up. ²⁴ All subjects had their blood pressure monitored in the office (where control was < 140/90 mm Hg) and outside the office (where control was < 135/85 mm Hg). There was no significant difference in prognosis between those whose blood pressures were controlled both at and outside the office (n = 3827, or 33.3% of the population studied) and those whose blood pressures were controlled only at home but not at the office (n = 1550, or 13.5% of the population studied). These observations support the concept that white coat hypertension has a better prognosis than sustained hypertension. The risk of individuals with masked hypertension (n = 1306, or 11.4% of the population studied: controlled blood pressure at the office but not at home) was not significantly different from that of sustained hypertensives in both locales (n = 4819, or 41.9% of the population studied), and both of these groups had a worse prognosis than that of those with controlled hypertension.

hypertension and a nonsignificant and only slightly higher risk than the 3827 with hypertension controlled in both the office and at home (Fig. 9-2). ²⁴ An intermediate viewpoint is that white coat hypertension is merely regression to the mean among the subset of patients with high BP variability.

Masked hypertension is said to be present when the in-office BPs childcare responsibilities, this condition is now found more widely, in perhaps 6% to 12% of the general population. ²⁴ As such patients would normally not be offered drug treatment (as their in-office BPs are, by definition, below treatment thresholds), masked hypertension is associated with a higher prevalence of target organ damage and incidence of CVD events than in true normotensives. In the seven published observational studies, the 1306 patients with this form of hypertension had only a slightly (and nonsignificant) reduced risk of future CVD events compared with the 4819 patients with sustained hypertension. ²⁴ Many have therefore recommended ABPM (or at least home readings) for all people at risk for

from prospective epidemiological surveys. Perhaps the most well

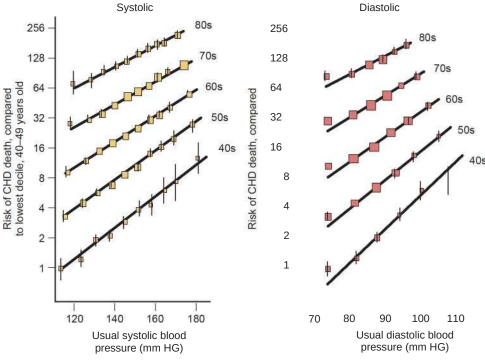


FIGURE 9-3 Relationship of risk of death from CHD (on a logarithmic scale) and the initial blood pressures (systolic at left, diastolic at right) measured at the beginning of each decade of life (given at the top of each line, from 40 to 89 years) in nearly 1 million participants in 61 epidemiologic studies. Larger boxes indicate estimates with smaller 95% confidence intervals, which are indicated for smaller boxes by the vertical lines. The total number of fatal CHD deaths was 30,143. The best-fit regression line for each decade of life ignores the point corresponding to the lowest blood pressure (always <115/70 mm Hg), which includes some individuals with *Very* low blood pressures. (Modified from Prospective Studies Collaborative. Lancet 360:1903, 2002.)

the Framingham Heart Study, in which 5209 healthy men and women were extensively evaluated initially and then observed over time. After a sufficient number of subjects had events, a quantitative estimate could be made of the importance of hypertension in the development of these events, even after adjusting statistically for the presence of other risk factors (eg, elevated plasma lipid levels or smoking). Data from Framingham and 60 other observational and epidemiologic data bases have been pooled (Fig. 9-3) and clearly show a strong, positive, and continuous relationship between the level of initial BP and the future risk of death from CHD. 12 Within each decade of life, for each BP increase of 20/10 mm Hg, beginning at 115/75 mm Hg, the risk of death from CHD, stroke, or CVD doubles. 12 These data also show that systolic BP is a much better predictor than diastolic BP or even pulse pressure of CVD and CHD outcomes. 12

Perhaps more important than epidemiological and observational studies of large numbers of people that correlate risk of death from heart disease and BP levels many years earlier are the results of prospective, randomized clinical trials that show, separately and in aggregate, that antihypertensive drug therapy reduces the risk of CHD and other CVD events during a < 6-year time frame. Most impressive are the results of studies comparing placebo or no treatment with antihypertensive drugs (typically diuretics and beta blockers in older studies, although a few studies with angiotensin-converting enzyme [ACE] inhibitors or calcium antagonists exist). Figure 9-4 shows the results of meta-analyses of 32 clinical trials that compared placebo or no treatment with an initial diuretic, beta blocker, calcium antagonist, or ACE inhibitor in the

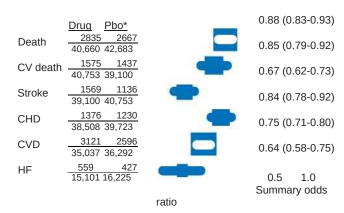


FIGURE 9-4 Results of meta-analyses of clinical trials involving placebo or no treatment compared with an *initial* low-dose diuretic, beta blocker, ACE inhibitor, or calcium antagonist in the prevention of cardiovascular events in hypertensive subjects. The results, all of which are highly significant (with no significant inhomogeneity), change only slightly if trials that gave the randomized agent as add-on therapy are included. The numbers above or below the line for each event are the numbers of subjects with that event or at risk for that event. *Pbo, placebo or no treatment; OR (95% CI), odds ratio (95% confidence interval); CV, cardiovascular; CHD, coronary heart disease; CVD, cardiovascular disease (first CHD event, stroke, or CV death); HF, heart failure. (*Updated from Elliott WJ: Cardiovascular events in clinical trials of antihypertensive drugs vs. placebo/no treatment: a meta-analysis [abstract*], J Hypertens 23[Suppl 2]:S273, 2005.)

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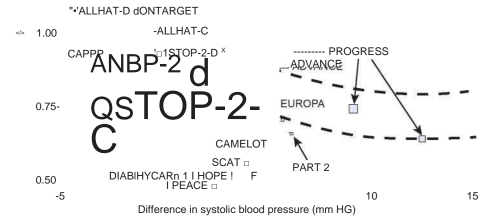


FIGURE 9-5 Meta-regression of the relative risk for CHD (*y*- axis, test agent-control) on the achieved difference in systolic blood pressure between randomized groups (control-test agent) in 27 clinical trials involving ACE inhibitors (placebo control in open squares, active control in filled squares). The regression line and its 95% confidence intervals are based on data from 27 older clinical trials. ²⁶ ADVANCE, Action in Diabetes and Vascular disease: preterAx and diamicroN-MR Controlled Evaluation; ALLHAT-D, Antihypertensive and Lipid-Lowering treatment to prevent Heart Attack Trial-Diuretic comparison; ALLHAT-C, Antihypertensive and Lipid-Lowering treatment to prevent Heart Attack Trial-Calcium antagonists comparison; ANBP-2, Australian National Blood Pressure trial 2; CAMELOT, Comparison of AMlodipine vs. Enalapril to Limit Occurrences of Thrombosis; CAPPP, CAPTOpril Primary Prevention Project; DIABHYCAR, DIABETES, Hypertension microalbuminuria or proteinuria, Cardiovascular events And Ramipril; EUROPE, EURopean Reduction Of cardiac events with Perindopril in stable coronary artery disease; HOPE, Heart Outcomes Prevention Evaluation; ONTARGET, ONgoing Telmisartan Alone and in combination with Ramipril Global Endpoint Trial; PART-2, Prevention of Atherosclerosis with Ramipril Trial 2; PEACE, Prevention of Events with Angiotensin-Converting Enzyme inhibition; PROGRESS, Perindopril Protection Against Recurrent Stroke Study; SCAT, Simvastatin/enalapril Coronary Atherosclerosis Trial; STOP-2-C, Swedish Trial in Old Patients with hypertension 2-Calcium antagonist comparison; STOP-2-D, Swedish Trial in Old Patients with hypertension 2-Diuretic or beta blocker comparison; UKPDS, United Kingdom Prospective Diabetes Study. (Updated from Elliott WJ, Jonsson MC, Black HR: Management of hypertension: is it the pressure or the drug? Circulation 113:2763, 2006.)

prevention of CVD in hypertensive subjects. These data indicate a highly significant benefit of BP-lowering drugs in hypertensive individuals across all types of CVD *and* all-cause mortality.

Perhaps the most direct and persuasive evidence in favor of the link between BP lowering and CVD (especially CHD) events in clinical trials can be shown in one of several meta regression analyzes correlating the difference in achieved systolic BP across randomized arms in large numbers of clinical trials involving hundreds of thousands of subjects with the relative risk for the specific CVD event (Fig. 9-5). 13,14,25,26 With the exception of heart failure, these analyzes generally show that a larger difference in achieved systolic BP (rather than the change in BP or initial BP) is associated with a larger difference between the randomized arms in CVD endpoints. For example, in the first such report, the differences in CHD across randomized treatment groups in trials was not at all well explained by the initial BPs of the participants in each trial (r = 0.02; P = 0.37) but was instead highly significantly correlated with differences across groups in achieved systolic BP ($r^2 = 0.53$; P = 0.0005). ²⁵ Such analyzes have also been used to claim "benefit beyond BP control" for specific antihypertensive drug classes in preventing specific end points (eg, ACE inhibitors preventing CHD), 14,27 but this is controversial 26

These data (both epidemiological and those derived from clinical trials) have been interpreted in several different ways. Several groups have pointed out that elevated BP is but one (relatively minor) predictor of CVD ²⁸; age, for example, is a much more powerful risk factor. The sensitivity and specificity of BP are low; not everyone with elevated BP will eventually have an event, just as, regrettably, not every person with a

"normal" BP will be spared. Any strategy that attempts to fix a value above which everyone should receive treatment is unlikely to be successful and cost-effective. These authorities recommend instead that treatment decisions about BP should be based on a person's absolute risk of CVD, which can be easily estimated by one of several country-specific risk estimators 28; the Framingham risk equation is the most widely used of these in the United States and has been recently updated with a risk algorithm for total CVD risk. ²⁹ For these reasons, their opinion about BP can be easily summarized: "Hypertension: Time to move on." Advocates of the polypill consider the results of BPlowering trials so compelling that universal treatment for everyone above a gender-specific age with three moderate-dose antihypertensive drugs (and other agents) is recommended, without even measuring BP. 15 These two different approaches lead to attempts to improve the cost-effectiveness of BP treatment. The first approach would provide treatment for individuals above a certain 10-year risk

among individuals with above-normal BPs. Hypertension

to see how JNC 8 deals with these suggestions.

PREDICTORS OF HYPERTENSION

In addition to demographic factors, many features of modern American life appear to increase the risk of becoming hypertensive. As shown in Figure 9-1, older age is probably the most powerful predictor. 8 For those younger than 55 years, men are at greater risk than women, but this reverses after the age of 55 years. 8 African Americans have a greater risk of hypertension as well as more target organ damage when it is diagnosed and a greater burden of heart disease, stroke, and end-stage renal disease, although these have been improving during the last 5 to 10 years. 6,8 Uncontrolled hypertension appears to be even more common in Mexican Americans than in other racial and ethnic groups, perhaps because of limited access to health care. 6,8 Although only a few clinical syn dromes are clearly hereditary, a single BP measurement is about 40% predictive of inherited BP levels; longitudinal measurements increase this to about 55%. 8 Prospective longitudinal studies of the heritability of BP in male medical students show about equal inheritance from the mother or father. 30

level (and those below it would fund their own medical care for

hypertension). The second approach uses only generically

available drugs (combined in a single pill) and avoids expensive

interactions with health care providers. It will be very interesting

Although lower educational attainment, socioeconomic status, lower physical activity scores, smoking, dyslipidemia, inflammation, psychosocial stressors, sleep apnea, and dietary factors (including higher consumption of dietary fats, sodium, alcohol, and calories or lower potassium intake) have all been implicated in many observational studies as significant predictors of hypertension, 8 much emphasis has recently been focused on obesity, particularly in childhood and adolescence. A study of the NHANES 1988-2004 data suggests that 20% to 80% of the increase in hypertension prevalence in adults during this period could be accounted for by the alarming increase in the average body mass index. 31 Many intervention trials have also demonstrated the benefit of a low-sodium diet in preventing hypertension, particularly one that is rich in fruits and vegetables and low in saturated fat. 32 Observational studies have also shown long-term benefit in preventing hypertension (and CVD events) of dietary pat terns similar to those in the Dietary Approaches to Stop Hypertension (DASH) cookbooks, ³³ which are among the most popular downloads from the website of the National Institutes of Health. 3.32

As with all diagnoses that depend on crossing a threshold of a continuous variable (eg, BP, fasting glucose concentration), probably the most important predictor of hypertension is prehypertension (BP 120-139/80-90 mm Hg), 2.3 simply because individuals with it are closer to BP > 140/90 mm Hg than are people with normal BPs (< 120/80 mm Hg). The TRial Of Prevention of HYpertension (TROPHY) was organized to test, in humans (as had been demonstrated in rodents), whether shortterm drug administration could prevent hypertension, even after the drug had been discontinued. ¹⁷ After 2 years of candesartan, the risk of hypertension was reduced by 66% compared with placebo (only three people needed treatment to prevent one from progressing); but after 2 years of receiving placebo, the benefit was much reduced (to 15.6%). Although adverse events were not significantly different, serious adverse events (including cardio vascular events: 1 versus 6) were less common in the group given candesartan. These data clearly showed that drug treatment of candesartan was feasible, but whether it is worth the cost is still debatable, particularly as a shorter Danish study showed no benefit after discontinuation of candesartan. 34

Another important predictor of hypertension is the use of nonsteroidal anti-inflammatory drugs (NSAIDs), particularly

Four key issues must be addressed during the initial evaluation of a person with elevated BP readings:

- Documenting an accurate diagnosis of hypertension. 18,23,28 9
- Stratifying the person's risk for CVD disease, which involves (1) defining the presence or absence of existing CVD or renal disease or target organ damage related to hypertension and (2) screening for other CVD risk factors that often accompany hypertension to obtain an esti mate of global cardiac risk. 3-5.9
- Assessing whether the person is likely to have an identifiable cause for the elevated BP (secondary hypertension) and should have further diagnostic testing for it. 3-5.9
- Obtaining information that may be helpful in choosing appropriate therapy. 3-5.9

Although extensive and expensive laboratory studies are rarely necessary in the evaluation of hypertensive patients, the health care provider must be able to recognize when additional studies or consultation with a specialist is appropriate and warranted. Delaying discovery of a potentially curable form of hypertension or failing to properly assess whether target organ damage or comorbidities are present puts the patient at unnecessary risk, delays implementation of specific treatment, prolongs the time to BP control, and increases CVD risk.

Documenting the Diagnosis

BP should be measured under relaxed and controlled conditions after appropriate rest (typically 5 minutes) and taken by someone whose ability to perform the measurement accurately has been certified. Because ~ 10% to 15% of Americans cannot properly hear and interpret Korotkoff sounds, it is unlikely that measurements reported by every observer are accurate. Excellent programs are available that can train, validate data, and recertify the competence of the person performing BP measurements. Careful attention should be given to proper, standardized technique. 18

Before the diagnosis of hypertension is made, an individual should usually have elevated BP measurements documented at least twice, at visits separated by a week or more. Each measurement should be an average of two or three readings differing by < 5 mm Hg from each other, taken a few minutes apart in the seated or supine position. Patients who exhibit wide fluctuations in BP or who are hypertensive at some evaluations but normotensive at others may need additional measurements, in the office, at home, or by ABPM, to confirm that they are indeed hypertensive. 3,18-23 Treatment should generally not be instituted until the diagnosis is clearly proven. In some circumstances, such as when target organ damage is present, treatment may need to be started after a single set of measurements.



¹²⁸ Stratifying Risk for Cardiovascular Disease

Before a treatment program directed at lowering BP is started, a thorough assessment of the person's risk for developing CVD is warranted. Little guidance is provided in JNC 7 about this, ³ but many other schemes are available to assist the physician, most of which have been developed for or adapted to European populations. ³⁶⁻³⁸ Many of these are semi-quantitative, similar to those of the Framingham Heart Study's risk score, ^{29,36-38} and allow a reasonable estimate of the E patient's 10-year risk of CVD to be calculated. The Sheffield

tables, based on the Scottish national health surveys, may be the most interesting of these. 38 These reasonably well dated tables (one for men, another for women) use age and 9 the ratio of total cholesterol to HDL-cholesterol (with 22 choices for men, 18 for women) and simply dichotomize hypertension (defined as > 140/90 mm Hg or taking treatment), smoking, and diabetes. Perhaps surprisingly, their data indicate that hypertension (whether treated or untreated) affects the overall risk of CHD only slightly; when other risk factors are incorporated into the risk equation, the absolute level of blood BP (especially at diagnosis) is almost irrelevant. 38

The determination of an individual's absolute risk has important implications for the selection of antihypertensive agents, the BP target, and the time to goal attainment. Individuals with the highest short-term risk of a stroke or heart attack are those who have already established CVD or renal disease (eg, history of a recent transient ischemic attack or previous myocardial infarction). These individuals should be treated promptly, intensively, and (in general, although controversy exists ³⁹) to a lower BP goal than for uncomplicated hypertensive people. ^{3-5,9,40,41} The search for evidence of concomitant CVD or renal disease need not be extensive or expensive, however. Typically, a complete medical history, a directed physical examination, and a few routine laboratory tests (including an electrocardiogram, urinalysis, and serum chemistry panel) are sufficient.

Hypertensive people with target organ damage are also at substantial risk for cardiovascular events. Target organ damage encompasses many subclinical features of the physical examination or laboratory test results indicating that there has already been an alteration of structure or function in the eyes, heart, kidneys, or blood vessels related to hypertension. Although such individuals may not have as yet suffered an irreversible hypertension-related event (eg, stroke), some are at substantial risk for these sequelae (eg, those with chronic kidney disease [CKD]), and the presence of target organ damage usually indicates that hypertension has been present for some time. These people also should receive prompt and intensive efforts to lower BP, typically to a lower than usual goal. Risk calculations are useful only for reasonably low-risk individuals. People with prior CVD, diabetes, or CKD are all assumed to have a 10-year risk of CVD > 20%. 3-5,9

Other CVD risk factors (tobacco use, family history of premature CVD) are often found in hypertensive people, and central obesity, dyslipidemia, diabetes, and hypertension tend to cluster. Because other risk factors tend to be additive (if not multiplicative) in increasing the probability of CVD events, it is important to screen a newly diagnosed hypertensive person for these other risk factors to more accurately estimate cardiovascular and renal risks.

Even though age is the most important (nonmodifiable) predictor of CVD risk (see Fig. 9-3), the treatment scheme and BP goals recommended in JNC 7 are independent of the patient's age. ³ There are now good data, several meta-analyses, ⁴¹ and an important recent placebo-controlled clinical trial ⁴² showing that older people, even past the age of 80 years, benefit greatly from BP-lowering drug therapy.

Considering Secondary Hypertension

More than 95% of Americans with hypertension have no specific cause of their elevated BPs (ie, idiopathic, essential, or primary hypertension). There are three reasons to consider the possibility that hypertension in a newly diagnosed patient might have a specific cause. First, BP control is often difficult to achieve in those with secondary causes of hypertension; diagnosis of it early is likely to get BP to goal more quickly. Second, and particularly important in younger people, use of specific modalities to cure the underlying disease will reduce the future burden of treatment (medical care costs, adverse effects of therapy, and quality of life). Last, routine consideration of secondary causes of hypertension when the diagnosis is originally made will ensure that these diagnostic possibilities—will be entertained, and the pros and cons of further testing critically evaluated, and that the clinician will not miss a secondary cause when it is present (see later for details).

Guided Therapy

The more than 120 antihypertensive agents and fixed-dose combinations currently available in the United States differ in BP-lowering efficacy in various situations. It is often helpful to discuss these potential confounders of treatment with the patient in an effort to "individualize" treatment according to the patient's specific dietary, medical, and personal considerations. For example, diuretics and calcium antagonists are more effective than ACE inhibitors and angiotensin II receptor antagonists when dietary sodium is excessive. JNC VI and JNC 7 both recommend treating hypertension and a concomitant illness or condition with a specific antihypertensive drug when that drug has been shown in clinical trials to improve CVD morbidity and mortality (so-called compelling indication for a specific class of antihypertensive drug). ³Therefore, even though ACE inhibitors were not routinely recommended as initial therapy for uncomplicated hypertensives, if the patient has heart failure of the systolic type, an ACE inhibitor could be prescribed. It would be expected not only to lower the BP but also to provide the impressive benefits seen in many long-term studies in every stage of heart failure. Last, some patients are particularly fearful of specific potential adverse effects of certain antihypertensive drugs (eg, male sexual dysfunction). If the prescriber knows this information, efforts may be taken to avoid medications with a high incidence of this particular problem.

Medical History

In addition to assessment of the risk of future CVD and renal disease, a careful medication, environmental, and nutritional history should be obtained during the initial evaluation and intermittently thereafter. It is particularly important to ascertain whether the patient is taking any agent (either by prescription or over-the-counter) or other substance that might elevate BP (Box 9-2). 43 Of particular concern are the NSAIDs, which are widely used and available over-the-counter and are sometimes not recognized as "medicines" by many patients. Sympathomimetic amines (once commonly found in weight loss, cold, and allergy preparations) have been associated with an increase in both BP and risk of intracerebral hemorrhage and stroke. Hypertensive people should avoid both NSAIDs and sympathomimetic amines and attempt to obtain relief of pain with acetaminophen and of the symptoms of nasal congestion with antihistamines, if possible. When these modalities are ineffective, short-term use of the usually prescribed drugs may be condoned, but with the recognition that BP control is



BOX 9-2 Substances That Can Raise Blood Pressur (Partial List)

Nonsteroidal anti-inflammatory drugs (including the newer COX-2 inhibitor celecoxib)

Corticosteroids

Sympathomimetic amines

Oral contraceptive hormones

Methylxanthines (including theophylline and caffeine*)

Cyclosporine

Erythropoietin

Cocaine

Nicotine 1 "

Phencyclidine (PCP)

COX-2, cyclooxygenase 2; PCP, phenylcyclohexyl piperidine ("angel dust").

Abridged from Elliott WJ: Drug interactions and drugs affecting blood pressure control. *J Clin Hypertens (Greenwich)* 8:731, 2006.

likely to be suboptimal during and immediately after their consumption.

Oral contraceptive pills containing estrogens and progestins may raise BP in some women, although this is much less of a problem with the lower doses in common use today. If a newly diagnosed hypertensive woman uses these pills, discontinuance for 6 months and observation of the BP may allow a decision to be made about whether the pills are the cause of hypertension. Conjugated estrogens (with or without progesterone), typically given for postmenopausal hormone replacement therapy, rarely raise BP, although they have been associated with a wide variety of other problems, including increased rates of CVD events.

Other prescription drugs can either elevate BP or interfere with certain antihypertensive agents. ⁴³ Of the former, cyclosporine, erythropoietin, corticosteroids, cocaine, and theophylline are perhaps the most widely recognized. Of the latter, monoamine oxidase inhibitors, NSAIDs, and tricyclic antidepressants are the most common. It is important to ascertain whether a hypertensive patient has taken any of these agents as well as several other illicit drugs (eg, phencyclidine). Some chemical elements, particularly lead and chromium, may elevate BP long after exposure; questioning about these and other environmental toxins may sometimes be helpful.

A focused dietary history is important because the most effective lifestyle modifications involve limiting either calories or sodium or both. 2-5,10,31,44,45 Dietary salt and saturated fat intake can be estimated from an informal survey of dietary habits and preferences. Many processed foods, "fast foods," "diet foods," condiments, and snack items are concentrated, oftenunrecognized sources of salt. Now that most foodstuffs bear labels attesting to their high sodium content, many patients are more easily able to choose healthier foods. A sensible target (now validated in several clinical trials, notably the DASH-Sodium substudy 46 and PREMIER 47) is 100 mEq (2.4 g or 2400 mg) of sodium per day; this can usually be achieved if the high-salt items mentioned before are avoided and the patient does not add salt either at the table or in cooking. On occasion, it is useful (and relatively inexpensive compared with formal dietary counseling) to have the patient collect a 24-hour urine sample for sodium, particularly when the patient claims to be avoiding salt but the physician is doubtful. Although not all hypertensive patients will experience a reduction in BP on a low-salt diet or an increase in BP on a high-salt diet, individuals who are "salt sensitive" (~60% of the US population) will benefit from reducing dietary sodium. In general, African Americans and elderly, obese, and diabetic patients are more likely to be salt sensitive, with BPs that are more responsive to dietary salt restriction.

The nutritional history should also include questions about

saturated fat consumption, dairy intake, and whether any mineral or vitamin supplements are being used. Because obesity is a major problem for many hypertensive patients, the calorie intake, eating pattern, and changes in weight should be included. Weight loss remains the most successful of all short-term lifestyle modifications for hypertension and should be a part of the therapeutic plan in all overweight hypertensives from the outset. 3,30,32,44

Social History

Although alcohol in moderation (one drink for women and two drinks for men maximum, with each drink being 12 ounces of beer, 4 ounces of wine, or 1 ounce of spirits) appears to protect against CHD in hypertensives, ⁴⁸ excessive alcohol intake (four or more drinks per day) raises both BP and all-cause mortality. In some patients, reducing or stopping alcohol ingestion can have salutary effects on BP. A clinical trial in veterans who consumed approximately six drinks per day when enrolled was unsuccessful in demonstrating a significant reduction in BP in those receiving a cognitive-behavioral alcohol reduction program, despite consuming significantly fewer drinks per day (by 1.3, on average) than the control group, who received a much less intensive educational program. Nevertheless, a meta-analysis suggests that there is a place for alcohol restriction in the non-pharmacological therapy for hypertension. ⁴⁹

Large populations of smokers have, on average, lower BP than nonsmokers do, probably because smokers tend to be less obese than nonsmokers. Consuming tobacco has both acute and chronic adverse effects on BP and hypertensive patients. Smoking a single cigarette raises BP and heart rate acutely (within seconds to minutes) because of nicotine's stimulation of catecholamine secretion. This effect disappears in about 15 minutes, so BP should be measured at least 15 to 30 minutes after the most recent cigarette is extinguished. Chronic tobacco abuse roughly doubles the long-term risk of CVD and has an even larger effect on peripheral arterial disease (including renovascular hypertension). Inquiry about tobacco abuse and advice to discontinue it (if present) should be a part of every encounter with a health care professional. Hypertensive patients should also be questioned about a sedentary lifestyle and whether there is willingness or ability to engage in regular physical activity. Even limited aerobic exercise, including brisk walking for 30 minutes on most days, can reduce BP and the risk of all-cause and CVD mortality. Snoring, daytime sleepiness, and other clinical features of obstructive sleep apnea, especially in obese hypertensives, should lead to a Berlin Questionnaire 50 and consideration of a formal evaluation for this underappreciated and underdiagnosed form of secondary hypertension. 51

Physical Examination

The directed physical examination of the hypertensive patient should pay special attention to weight, target organ damage, and features consistent with secondary hypertension . It should focus on items that were suggested by the medical history.

The pattern of fat distribution should be noted. Android obesity (waist-to-hip ratio > 0.95) is associated with increased CVD risk, whereas gynecoid obesity (waist-to-hip ratio < 0.85) is not. Men whose waist is > 102 cm (40 inches) and women whose waist is > 88 cm (35 inches) are at increased risk. The International Diabetes Federation recommends lower waist circumference cut points



^{*}Short duration (minutes to hours).

¹ Very short duration (seconds to minutes).

- for other populations, such as European whites and Asians, to define abdominal obesity.
- The patient's skin should be carefully examined for cafe au lait spots (suggesting neurofibromatosis and possible pheochromocytoma), acanthosis nigricans (suggesting insulin resistance), xanthomas at tendons, or xanthe lasma (indicating dyslipidemia). Other skin signs suggestive of pheochromocytoma (axillary freckles, ash-leaf patches, port-wine stains in the trigeminal distribution, and adenoma sebaceum) are uncommon, except in patients with phakomatoses.
- The many physical signs associated with other secondary causes should be sought, particularly if they are suggested by the medical history. The signs of Cushing syndrome (purple striae, moon facies, dorsocervical fat pad, atrophic skin changes) or thyroid disease (abnormal Achilles reflexes, hair quality, and eye signs) are typically difficult to ignore.
- The funduscopic examination is important in assessing the duration and severity of hypertension. The presence of hypertensive retinopathy (grade 1: arterial tortuosity, silver wiring; grade 2: arteriovenous crossing changes ["nicking"]; grade 3: hemorrhages or exudates; grade 4: papilledema) provides definitive evidence of target organ damage.
- The neck should be examined for an enlarged thyroid gland, abnormalities of the venous circulation (eg, jugular venous distension, abnormal or cannon *a* waves), and carotid bruits.
- The chest should be auscultated for evidence of heart failure or bronchospasm; bronchospasm would likely make beta blockers contraindicated.
- The heart should be examined carefully for cardiomegaly, murmurs, and extra sounds.
- The abdominal examination is one of the most important parts of the directed physical examination because the finding of an abdominal bruit is one of the most cost-effective ways to screen for renovascular hypertension. All four abdominal quadrants and the back should be auscultated, typically with the use of the pulse at the wrist as the synchronizing stimulus. Diastolic or continuous bruits are common in renovascular hypertension, but systolic bruits in young and especially thin hypertensive subjects may not be indicative of renal artery stenosis. Abdominal masses can sometimes be palpated in patients with pheochromocytoma or poly cystic kidney disease.
- The groin and legs should be examined for evidence of peripheral arterial disease, which is often manifested as bruits, absent or decreased pulses, and abnormal hair growth patterns. Edema can be a sign of heart failure, renal disease, or high doses of dihydropyridine calcium antagonists.
- The neurological examination need not be extensive in a hypertensive patient with no history of cerebrovascular disease, but it should be complete if a history of stroke or transient ischemic attack is present.

Laboratory Testing

For most hypertensives, only a few simple and inexpensive laboratory tests are needed initially. In selected patients, however, more extensive testing is not only appropriate but also necessary to diagnose secondary hypertension and to avoid delaying proper treatment. The laboratory tests that are recommended for all hypertensive persons are shown in Box 9-3 and can be divided into those that are done to assess risk, to

establish etiology, to screen for important common diseases, and, finally, to guide the choice of initial therapy.

BOX 9-3 Laboratory Tests Appropriate for All Newly Diagnosed Hypertensive Patients

Assessing Risk

Lipid profile (including serum total cholesterol, HDL-cholesterol, and triglycerides)

Serum glucose concentration (preferably fasting)

Serum creatinine concentration
Urinalysis (both dipstick and microscopic)
12-Lead electrocardiogram

Establishing Cause

Serum potassium concentration

Serum creatinine concentration

Urinalysis (both dipstick and microscopic)

Thyroid-stimulating hormone (with further testing, if abnormal)

Screening for Common Asymptomatic Diseases

Complete blood count
Serum calcium concentration

Guided Therapy

Lipid profile (including total cholesterol, HDL-cholesterol, and triglycerides) Serum glucose concentration (preferably fasting)

Serum creatinine concentration

For uncomplicated hypertensive patients whose history and physical examination do not suggest a secondary cause, the simple battery of tests in Box 9-3 is all that is needed. A lipid profile and fasting glucose concentration are indicated in hypertensive patients because of the high prevalence of the metabolic syndrome in hypertensives. The presence of diabetes or additional risk factors not only requires institution of therapy for these conditions but also indicates substantially increased CVD risk, thus requiring more intensive therapy for hypertension, a lower goal BP, and closer follow-up.

Routine measurement of serum creatinine and a full urine analysis are recommended for three reasons. First, the urinary test is useful to assess risk because hypertensive patients with microalbuminuria, proteinuria, or renal impairment have a distinctly worse prognosis. Second, the urinalysis may identify CKD as a secondary cause. Renal impairment (typically manifested as elevated serum creatinine) with a normal urinary sediment may be a valuable clue suggesting ischemic nephropathy, perhaps due to renal artery stenosis. Third, knowledge of the serum creatinine level frequently guides therapy ⁵² because loop diuretics are routinely needed and are more effective than thiazide or thiazide-like diuretics when the estimated glomerular filtration rate (eGFR) is < 30 mL/ min/1.73 m².

An electrocardiogram may provide important but limited information in most hypertensive patients. Although it may identify an occasionally important dysrhythmia or even an unsuspected old myocardial infarction, its biggest role is in screening for left ventricular hypertrophy, which is an objective measure of both the severity and duration of elevated BP. Despite a sensitivity of only 10% to 50% (depending on which criteria are used in its interpretation) in the Framingham Heart Study, electrocardiographic evidence of left ven tricular hypertrophy

was associated with an approximately threefold increase in incidence of CVD events. Left ventricular hypertrophy detected by echocardiography appears to be an even better predictor of future events, but this test is not

recommended for routine evaluation because of its high cost and high intrinsic variability of a single procedure (-10 % to 15%).

Because hypothyroidism is a cause of remediable hypertension that is subtle, especially in the elderly, a thyroid-stimulating hormone assay may be helpful. Serum calcium is useful in evaluating hyperparathyroidism and is often included in automated chemistry panels. The measurement of plasma renin activity is useful (with a plasma aldosterone concentration) in the diagnosis of mineralocorticoid excess states, such as primary hyperaldosteronism.

SECONDARY HYPERTENSION

Important clues to the presence of secondary hypertension are often provided by a carefully obtained medical history (see earlier). Many patients with primary hypertension report an isolated elevated BP reading some time in their 20s and 30s that was not reproducible or sustained until at least a decade or more later. The level of BP gradually rises until it reaches a threshold level, and then hypertension is diagnosed. In contrast, patients with an identifiable secondary cause of hypertension usually present with a very different history. Instead of the gradual onset of elevated BPs, they usually have a relatively abrupt onset of hypertension, typically presenting at a higher stage and with considerable target organ damage. The sudden onset of elevated BPs at ages before 30 years or after 50 years should alert the clinician to the possibility that the patient may have secondary hypertension . Thus, the history of the patient's presentation with hypertension should be carefully documented. At what age were the BP readings first elevated? How high were the reading scores? Were all previous readings within the normal range? Was it discovered during a routine office visit, or did the patient have clinical problems related to BP elevation or related target organ damage?

Because patients with secondary hypertension typically do not respond as well as patients with primary hypertension to antihypertensive drug therapy, the history of the patient's response to treatment must be ascertained. What drugs were used and at which doses? Did the patient's BP respond initially and then become resistant? A positive answer to this question is frequently found in patients with primary hypertension who later develop secondary hypertension, particularly atherosclerotic renovascular disease.

Laboratory Testing for Secondary Hypertension

All types of secondary hypertension are uncommon in the general hypertensive population, although certain factors (eg, obesity and snoring for obstructive sleep apnea, smoking and peripheral vascular disease for renovascular hypertension) make specific diagnoses more likely. In interpreting test results in a given patient, Bayesian analysis (which incorporates the pretest probability of finding disease) is therefore more important than the sensitivity of the test (ie, the percentage of people with disease who have a positive test result). Those tests that are often recommended for the evaluation of patients for secondary hypertension are listed in Table 9-3. The proper evaluation of patients for many forms of secondary hypertension is still highly controversial despite the recent publication of guidelines for each. 53-56 Probably the most controversial are those pertaining to renovascular hypertension, for which the popular angioplasty and stenting have yet to show a significant long-term benefit in the four trials completed to date. 57-59

TABLE 9-3	Screening and Other Tests for Secondary Hypertension	Common Forms of
Diagnosis Renovascular	Preferred Screening Tests	Other Tests
hypertension	Doppler ultrasound of rena arteries, computed tomographic angiography	l Magnetic resonance angiography, renal angiography, captopril scintigraphy
Mineralocorticoid excess states	Plasma aldosterone/ renin ratio, 24-hour urinary aldosterone during salt loading	Computed tomographic scan of adrenals
Pheochromocytoma 2	24-hour urine for vanillylmandelic acid and metanephrines	3
		Plasma metanephrines, plasma catecholamines, T2-weighted magnetic resonance imaging
Sleep apnea	Berlin Questionnaire	
		Formal home nocturnal somnographic study
Cushing syndrome	8 AM plasma cortisol level	Dexamethasone suppression tests
Hypothyroidism	Thyroid-stimulating hormone	Serum thyroxine, triiodothyronine levels
Modified from recent	reviews and consensus	53-57

BRIEF REVIEW: OUTCOMES-BASED CLINICAL TRIALS

Because so much information about hypertension and its treatment has been derived from clinical trials, many excellent, evidence-based recommendations can now be made about its diagnosis and management. ^{3-5,9} Although details of many of the older studies are available in an excellent monograph, ⁶⁰ the results of most of these trials have been combined and summarized in other forms (eg, meta-analyses ^{4,13-15,24-27,36,41,45}), which, when properly done, are said to rank higher than individual trials in the hierarchy of evidence-based medicine.

Diuretics and Beta Blockers

In the early 1990s, the most frequently cited meta-analysis compared the observed results of BP lowering in clinical trials with those expected from epidemiological studies. Although stroke reduction in clinical trials (by about 42% ± 6%) was almost exactly what would have been expected, prevention of CHD was substantially less (about $14\% \pm 5\%$ compared with the expected 25% to 28%). 61 This led some to believe that the drugs used in the early trials (almost exclusively diuretics and beta blockers) might be somewhat less effective in preventing CHD than stroke. A subsequent network meta-analysis of these and later trials showed that an initial diuretic or beta blocker was associated with a higher risk of incident diabetes. 62 Similarly, an initial beta blocker (particularly atenolol) has been found in meta-analyses to be significantly less effective than any other initial antihypertensive drug in preventing most types of CVD events, 63 so this class has been demoted to fourth-line therapy in the most recent British guidelines. 4

Many of the placebo-controlled clinical trials of ACE inhibitors that had impressive results were performed in "high-risk

patients," not all of whom had hypertension, in which the randomized agents were added to whatever other antihypertensive and other drugs were appropriate. Some therefore object to the inclusion of these studies in meta-analyses that attempt to summarize the results of BP-lowering therapies.

Rather than asserting, for example, that ramipril is a better ACE inhibitor than trandolapril for prevention of CVD events, such trials can be used to illustrate the importance of concurrent therapies and residual absolute risk. In the Heart Outcomes Prevention Evaluation (HOPE), 64 only about 28%, 39%, and 76% of the patients were taking lipid-lowering agents, beta blockers, or aspirin (respectively), and the absolute risk of the primary outcome in the placebo-treated group was 39.4 events/1000

patient-years of follow-up. In the Prevention of Events with Angiotensin-Converting Enzyme inhibition (PEACE) trial, the drug use rates were 70%, 60%, and 90% (respectively), so the absolute risk with placebo was reduced to only 21.1 events/1000 patient-years of follow-up. 65 It was therefore much more likely that the HOPE investigators would see a significant benefit of

ramipril compared with trandolapril in PEACE.

Several meta-analyses and meta-regressions have suggested that ACE inhibitors have a specific protective effect on CHD events, independent of BP lowering, 14,27 but this is controversial ²⁶ The most recent trial involving an ACE inhibitor, the ONgoing Telmisartan Alone and in combination with Ramipril Global Endpoint Trial (ONTARGET), 66 compared ramipril and telmisartan (and their combination) and showed no evidence of "benefit beyond BP control" with the ACE inhibitor (which was given along with 1.18 other antihypertensive drugs, on average). Importantly, ONTARGET also showed no significant CVD event benefit with the addition of telmisartan to ramipril, despite a slightly lower BP in subjects randomized to the combination. placebo-controlled trials were disappointing because these

Calcium Antagonists

Although fears were raised in the mid-1990s by case-control, cohort, and meta-analytic studies that calcium antagonists were associated with a significantly higher risk of CHD, many subsequent randomized clinical trials, especially Antihypertensive and Lipid-Lowering treatment to prevent Heart Attack Trial (ALLHAT) ⁶⁷ and the Anglo-Scandinavian Cardiac Outcomes Trial (ASCOT), ⁶⁸ showed a slightly *lower* risk of CHD with amlodipine than its comparators. Furthermore, an initial calcium antagonist appears to be slightly (but not significantly) better than an initial diuretic in preventing a first stroke in hypertensive patients, ^{13,14,25-27,41} again allegedly independent of BP control. ²⁷ Some attribute the benefits seen in ASCOT to the lower central aortic pressure with the amlodipine-based regimen compared with the atenolol-based regimen. 69 Conversely, both dihydropyridine and nondihydropyridine calcium antagonists are treatment in all 55 published clinical trials that enrolled only hypertensive subjects (n = 261,051). associated with a higher risk of heart failure. 13,14,25-27,41

Angiotensin II Receptor Blockers

The evidence base for ARBs is limited, primarily because ethics subjects (13 trials, 11,390 events, 96,332 subjects) are included in similar meta-analyses or if prevents unconfounded comparison of this newer class of Bayesian meta-analyses are performed. ARB, angiotensin receptor blocker; ACE-I, angiotensin antihypertensive drugs with placebo in outcome studies. The study converting enzyme inhibitor; CCB, calcium channel blocker (or calcium antagonist); w, designs have been limited to specific indications (eg, type 2 diabetic incoherence value, reflecting how well the data "hang together" (lower values are better). nephropathy or heart failure), comparisons against other BP- (Updated from Elliott WJ, Basu S, Meier PM: Dihydropyridine vs. nonlowering drugs, or add-on therapy in high-risk patients (not all of dihydropyridine calcium antagonists as initial therapy for prevention of whom were hypertensive). 66,70,71 Although the ARB was cardiovascular events in hypertensive patients [abstract]. J Clin Hypertens "noninferior" to the ACE inhibitor in ONTARGET, 66 the (Greenwich) 11[SupplA]:A13, 2009.) nonsignificant results of the two

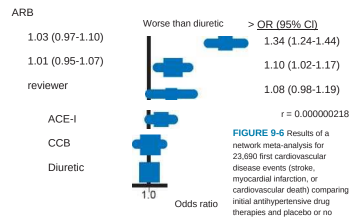
drugs are effective in lowering BP and are well tolerated.

Combination Therapy

Although nearly all clinical trials end up comparing drug regimens of multiple antihypertensive agents, two recent trials have provided data on the relative effectiveness of combination therapy. In ONTARGET, the combination of full -dose ramipril and full-dose telmisartan resulted in more BP lowering but significantly more adverse effects and renal end points than with ramipril alone. 66,72 The design of the Avoiding Cardiovascular events through COMbination therapy in Patients LIving with Systolic Hypertension (ACCOMPLISH) trial 73 was unique in that it set out to compare outcomes in a very high risk hypertensive population that was randomized to benazepril plus either hydrochlorothiazide (as recommended in JNC 7) 3 or amlodipine (as in the recent British guidelines). 4 The BP lowering in both groups was excellent, but the BP lowering was slightly better and CVD outcomes were less common in the group randomized to benazepril plus amlodipine. Unanswered questions about ACCOM PLISH include whether the chosen diuretic or the ACE inhibitor was optimal in dose and identity and what other explanation might be offered for the observed difference in outcomes. This trial has the potential to change the standard recommendation of JNC 7 regarding initial treatment of stage 2 hypertension, for which a diuretic and another antihypertensive drug were advised. 3

Summary of Current Evidence from Clinical **Trials**

There is little doubt, on the basis of the clinical trial evidence gathered to date and summarized before 4,13 · 15,25 · 27,41,63 and updated in Figure 9-6, 74 that BP lowering is a powerful means of achieving reductions in CVD events. The initial assessment



For some trials (eg, ALLHAT), the numbers of subjects experiencing a first event were estimated, as these data have not yet been reported. These odds ratios (OR, listed on the right, with 95% confidence intervals, 95% CI) change little (but in the expected direction) if other trials that compared add-on drug therapy (3 trials, 1889 events, 25,194 subjects) or enrolled nonhypertensive

of CVD risk takes on greater importance, as specific therapies are said to be more protective for different events: a diuretic to prevent heart failure, a calcium antagonist to prevent stroke, 15,27 and an ACE inhibitor to prevent CHD. 14,27 Multiple economic analyzes also indicate that in addition to being effective in preventing CVD events, antihypertensive drug therapy is also relatively cost-effective compared with other commonly employed treatment strategies used in medicine today. 4

MANAGEMENT

Successful management of hypertension requires a major commitment from the patient, the health care provider, and the health care system. The patient must continue to take what could be a costly medication with potential adverse effects and see a health care provider frequently, for an asymptomatic condition, in optimal drug choice. 3-5 the belief that this will reduce the risk of a major complication or even death. The physician must help the patient achieve BP goal and maintain surveillance on this and other CVD risk factors, without being sure that such treatment will prevent an event that Since the early termination of the alpha blocker (doxazosin) arm would have occurred without it. The health care system must fund and the final results of ALLHAT, the "default choice" for initial both pharmacy and provider benefits, understanding that the 5- antihypertensive drug therapy for most uncomplicated patients year number needed to treat (to prevent one expensive CVD event) in the United States is a low-dose thiazide-type diuretic, which is higher than its actuaries would prefer.

Lifestyle Modifications

measures to control BP. In the short term, weight loss is the most but it did recommend a thiazide-type diuretic "for most patients." effective, 44 but dietary sodium restriction is the only modality Some day, pharmacogenetic testing may be used to predict both shown to have a long-term benefit in reducing CVD morbidity and the BP-lowering response and the preventive effect of the mortality. 45 In the Trials of Hypertension Prevention, the difference thiazide-like diuretic, 82 but these techniques are currently in daily sodium excretion during the 18 to 48 months of follow-up unavailable to the general population. In the United Kingdom, between the randomized groups was only 816 to 1081 mg/day, 45 suggesting that the 2300 mg/day target recommended by inhibitor (or occasionally an ARB) for younger white patients, a guidelines and the DASH-Sodium and PREMIER trials is a good calcium antagonist for older or black patients, or a diuretic for one. 46,47 Also recommended (if appropriate for the individual) are those at high risk of heart failure. 4 The European Society of alcohol restriction to one or two drinks a day (two or three in the Cardiology/European Society of Hypertension guidelines recent British guide lines), tobacco avoidance, major exercise (30 recognize that few patients will achieve BP control with a single minutes most days of the week), reduction of caffeine (if excessive), agent, and therefore less importance is placed on which drug is and supplements of potassium, calcium, or magnesium (if and only chosen first. 5 The 2009 Canadian Hypertension Education if a deficiency state is present). There are few long-term controlled Program guidelines stratify even further on the basis of whether trials of yoga, ⁷⁵ paced breathing, ⁷⁶ spinal manipulation, ⁷⁷ fish oil, the BP elevation is systolic alone or diastolic ± systolic. ⁸³ For the ⁷⁸ or acupuncture ⁷⁹ for lowering of BP, although each can be useful former, a thiazide diuretic, long-acting calcium antagonist, or for selected patients. There is limited clinical trial evidence but ARB is recommended, but the diuretic and calcium antagonist good public health reasons for advocating a combination of weight have the best-quality evidence. For the latter, a thiazide diuretic, loss, dietary salt restriction, and other lifestyle modification beta blocker (for patients < 60 years of age), ACE inhibitor, or methods as preventive, 80 adjunctive, and (occasionally) definitive long-acting calcium antagonist is recommended, with the treatment of hypertension. 32,44-49 Recidivism can be a problem for thiazide having the best-quality evidence. 83 both weight loss and sodium restriction, however, and long-term adherence to such programs is uncommon in many clinics. The 20/10 mm Hg in BP with a single drug is low, many only study to assess outcomes using a vigorous program of lifestyle hypertension guidelines recommend consideration of initiating modifications with and without antihypertensive drug therapy with a two-drug combination for individuals with stage showed significant prevention of CVD events with the 2 hypertension (or a BP > 150/90 mm Hg for diabetics 3.84 or those combination. 81 As a result, for patients with comor bidities, target with either CKD 3.85 or established heart disease 86). JNC 7 organ damage, or high 10-year risk scores, many clinicians prefer suggested that a diuretic be part of such a regimen, but the to recommend lifestyle modifications plus appropriate drug ACCOMPLISH data call this strategy into question. 73 Few therapy.

Choice of Initial Drug Therapy

Initial Antihypertensive Drug Therapy for "Complicated Patients"

Since 1997, US hypertension guidelines have formally recognized that many people begin treatment too late, after they

have already developed CVD or other medical conditions that 133 may be positively affected by specific antihypertensive drug

therapies. ³ These have been divided into "compelling indications" and "clinical conditions." In the compelling indications (Table 9-4), a specific type of antihypertensive drug should be prescribed if clinical trials have demonstrated that class of drug to reduce morbidity or mortality for that condition. ³ Thus, for a hypertensive person with systolic heart failure, an ACE inhibitor will likely not only lower BP but also reduce CVD events and hospitalizations. There are also some absolute and relative contraindications for specific antihypertensive drugs that limit their use as initial therapy in all hypertensives (Table 9-5). Two examples of clinical conditions for which specific antihypertensive drugs benefit certain patients are a thiazide diuretic for calcium- g containing renal stones and a beta blocker for essential tremor. Although neither drug is likely to reduce CVD morbidity or mortality from these conditions, each will reduce symptoms and therefore improve adherence and BP control in the long term. JNC 7 and other hypertension treatment guidelines recognize the merits of specific antihypertensive drugs in complicated hypertensive patients, for whom an initial diuretic is sometimes not the

Initial Antihypertensive Drug Therapy for "Uncomplicated Hypertensives"

showed superiority in the secondary end point of combined CVD. Those who strictly adhere to evidence-based medicine will insist on chlorthalidone. 52 Perhaps because the 12.5-mg dose of chlorthalidone is not commercially available, JNC 7 did not All hypertension guidelines recommend nutritional-hygienic distinguish between chlorthalidone and hydrochlorothiazide, recommended initial antihypertensive therapy is usually an ACE

> Because the probability of achieving a reduction of more than subjects in ACCOMPLISH were "uncomplicated stage 2 hypertensives," however. 73

Compelling Indication	Treatment Prevents or Delays	Recommended in 1997	Recommended in 2003	Recommended in 2010
Heart failure (systolic type)	CV events	ACE-I (CONSENSUS, SAVE)	Beta blockers (MERIT-HF); spironolactone (RALES); ARB (Val-HeFT)	ARB (CHARM-Added)
After recent MI	Recurrent infarction or death	Beta blocker (ISIS)		
Diminished left ventricular functi after recent MI	ion Recurrent infarction, CHF hospitalization	ACE-I (SAVE, TRACE)	Eplerenone (EPHESUS)	ARB (VALIANT)
High risk for CVD	CVD events		ACE-I (HOPE)	
				ARB (TRANSCEND, ONTARGE
Type 1 diabetes mellitus	Deterioration in renal function	ACE-I (CCSG)		
Type 2 diabetes	CV events		ACE-I (MICRO-HOPE)	Diuretic + ACE-I (ADVANCE)
	Deterioration in renal function		ARBs (IDNT, RENAAL)	
Type 2 diabetic nephropathy				
Type 2 diabetes	Progression of microalbuminuria		ACE-I (MICRO-HOPE); ARB (IRMA- 2)	Diuretic + ACE-I (ADVANCE)
Older hypertensive persons	CV events	Diuretic (SHEP); DHP-CA (Syst-Eur)	ACE-I or DHP-CA (STOP-2); DHP-CA (Syst-China); ARB (SCOPE, second-line); ARB (LIFE)	Diuretic (HYVET)
	Deterioration in renal function		ACE-I (REIN, AIPRI, AASK)	
Nondiabetic renal impairment				
Prior stroke, TIA	Stroke and CV events		ACE-I (PROGRESS)	ARB (ACCESS, MOSES)
Left ventricular hypertrophy (usi strict criteria)	ng CV events (perhaps limited to stroke	?)	ARB (LIFE)	

ACE-I, angiotensin-converting enzyme inhibitor; ARB, angiotensin receptor blocker; CHF, congestive heart failure; CV, cardiovascular; CVD, cardiovascular disease; DHP-CA, dihydropyridine calcium antagonist; MI, myocardial infarction; TIA, transient ischemic attack.

CONSENSUS, COoperative North Scandinavian ENalapril SURvival Study (N Engl J Med 316:1429, 1987); SAVE, Survival And Ventricular Enlargement study (N Engl J Med 327:669, 1992); CCSG, Captopril Cooperative Study Group (N Engl J Med 323:1456, 1993); SHEP, Systolic Hypertension in the Elderly Program (JAMA 265:3255, 1991); Syst-Eur, Systolic Hypertension in Europe trial (Lancet 360:757, 1997); MERIT-HF, MEtoprolol Randomized Intervention Trial in congestive Heart Failure (JAMA 283:1295, 2000); RALES, Randomized Aldactone Evaluation Study (N Engl J Med 341:709, 1999); Val-HeFT, Valsartan Heart Failure Trial (N Engl J Med 345:1667, 2001); CHARM-Added, Candesartan in Heart failure Assessment of Reduction in Mortality and morbidity (Lancet 362:767, 2003); ISIS, International Study of Infarct Survival (Lancet 2:57, 1986); TRACE, TRandolapril Cardiac Evaluation (N Engl J Med 333:1670-1676, 1995); EPHESUS = Eplerenone Post-myocardial infarction Heart Failure Efficacy and Survival Study (N Engl J Med 348:1309, 2003); VALIANT, VALsartan In Acute Myocardial Infarction (N Engl J Med 349:1893, 2003); HOPE, Heart Outcomes Prevention Evaluation (N Engl J Med 342:145, 2000); ONTARGET, ONgoing Telmisartan Alone and in combination with Ramipril Global Endpoint Trial (N Engl J Med 358:1547, 2008); TRANSCEND, Telmisartan Randomized Assessment Study in ACE intolerant subjects with cardiovascular disease (Lancet 371:1174, 2008): MICRO-HOPE, MIcroalbuminuria, Cardiovascular and Renal Outcomes substudy of the Heart Outcomes Prevention Evaluation (Lancet 355:253, 2000); ADVANCE, Action in Diabetes and Vascular disease: preterAx and diamicroN- mr Controlled Evaluation (Lancet 370:829, 2007); IDNT, Irbesartan Diabetic Nephropathy Trial (N Engl J Med 345:841, 2001); RENAAL, Reduction of Endpoints in Non-insulin dependent diabetes mellitus with the Angiotensin II Antagonist Losartan (N Engl J Med 345:861, 2001); IRMA-2, IRbesartan MicroAlbuminuria study 2 (N Engl J Med 345:870, 2001); STOP-2, Swedish Trial in Old Patients with hypertension 2 (Lancet 354:1751, 1999); Syst-China, Systolic Hypertension in China trial (J Hypertens 16:1823, 1998); SCOPE, Study on Cognition and Prognosis in the Elderly (J Hypertens 21:875, 2003); LIFE, Losartan Intervention For Endpoint Reduction (Lancet 359:995, 2002); HYVET, Hypertension in the Very Elderly Trial (N Engl J Med 358:1887, 2008); REIN, Ramipril Evaluation In Nephropathy trial (Lancet 352:1252, 1998); AIPRI, Angiotensin-converting-enzyme Inhibition in Progressive Renal Insufficiency (Kidney IntSuppl 63:S63, 1997); AASK, African American Study of Kidney disease and hypertension (JAMA 288:2421, 2002); PROGRESS, Perindopril Protection Against Recurrent Stroke Study (Lancet 358:1033, 2001); ACCESS, Acute Candesartan CilExetil therapy in Stroke Survivors (Stroke 34:1699, 2003); MOSES, Morbidity and mortality after Stroke, Eprosartan compared with nitrendipine for secondary prevention (Stroke 36:1218, 2005).

TABLE 9—5 Contraindications for Specific Antihypertensive Drug Classes				
Antihypertensive Drug Class	Contraindications			
Thiazide diuretic	Allergy			
Beta blocker	Asthma, allergies			
ACE inhibitor	Angioedema due to ACE inhibitor, pregnancy, renal artery stenosis			
Calcium antagonist	Allergy			
Angiotensin II receptor blocker	Pregnancy, renal artery stenosis			
Alpha blocker	Orthostatic hypotension with frequent falls			
Alpha 2- agonist (centrally acting drug)	Allergy			

Add-on Drug Therapy

Perhaps because few clinical trials directly compared second-step antihypertensive drugs (after an initial diuretic), JNC 7 recommended an ACE inhibitor, ARB, beta blocker, or calcium antagonist but gave little further guidance. ³ Some would add an alpha blocker to the list, which was admittedly inferior to chlorthalidone as initial therapy in ALLHAT. 86 Others would routinely favor an ACE inhibitor or an ARB because of the potential synergy on serum potassium in combination with a thiazide or thiazide-like diuretic. Such a strategy has been more effective in achieving BP control during 6 months compared with traditional Canadian guidelines . 87 Beta blockers have the largest clinical trial experience as the second-line agent after a diuretic, but some object to the higher risk of incident diabetes with the combination. 4,62,68,84 Although the first published factorial-design trial of antihypertensive agents included a diuretic and calcium

INvestigation of Cardiovascular (CONVINCE) trial, 88 the most direct evidence in its favor is a was 53.5%, despite a mean BP of about 133/78 mm Hg after Chinese trial showing significant reduction in CVD (mostly stroke) completion of the randomized trial. 97 The < 130/80 mm Hg target events, comparing placebo against a calcium antagonist as second- for patients with CKD can be viewed as a reasonable compromise line therapy after a diuretic. 89

Target Blood Pressures

JNC 7 recommended only one of two BP targets. For uncomplicated hypertensives, the BP should be lowered to < 140/90 mm Hg. The best clinical trial evidence in favor of this target comes from the Hypertension Optimal Treatment (HOT) study. 90 It was the largest trial to compare outcomes in subjects randomized to different 39 Although the idea that these high-risk patients should be treated target BPs. Its 18,790 hypertensive patients achieved "optimal reduction in cardio vascular risk" with an achieved BP of 138.5/82.6 mm Hg. There was no significant increase in risk by further trial evidence comes from a post hoc analysis of the progression lowering BP to a diastolic target of < 80 mm Hg, but there was no of atherosclerosis in the Comparison of AMlodipine vs. Enalapril further benefit either. 90 An open-label study of 1111 Italian to Limit Occurrences of Thrombosis (CAMELOT) study. 98 nondiabetic hypertensives has recently suggested that systemic BP be lowered to < 130 mm Hg because it reduces both 1991 CHD patients with controlled BP (baseline, 129/78 mm Hg) echocardiographic left ventricular hypertrophy and cardio - randomized to additional treatment with amlodipine, enalapril, vascular outcomes. 91 However, because the probability of drug- or placebo, 99 a substudy of 274 patients used intravascular related side effects often increases as the doses are increased, it is ultrasound to measure atherosclerosis progression or regression likely that the target BP of < 140/90 mm Hg for uncomplicated . Individuals with on-treatment BP $\stackrel{•}{>}$ 140/90 mm Hg had a hypertensive patients will be defensible for some time to come.

BP target of < 130/80 mm Hg, ³ which is easier to justify for diabetic with BP < 120/70 mm Hg had a decrease in atheroma volume. ⁹⁸ patients (although both are controversial). 39 The first clinical trial Some are still concerned, however, that lowering of BP "too far" to demonstrate the benefit of a lower BP target for diabetics was the in patients with CHD can be deleterious. 100 The most recent data United Kingdom Prospective Diabetes Study (UKPDS 38). 92 to support the existence of the J-shaped curve comes from a post During 8.4 years of follow-up, 1148 type 2 diabetics randomized to hoc analysis of the International VErapamil/trandolapril STudy two different BP goals (<150/85 mm $^{\circ}$ Hg or <180/105 mm Hg) did (INVEST). 101 Although there were no significant differences in much better if treated to the lower goal. The achieved BP difference outcomes among CHD patients randomized to either atenolol or between groups was 10/5 mm Hg (154/87 versus 144/82 mm Hg), verapamil, CHD events were significantly more common in which if subtracted from the usual then-current BP goal for those whose diastolic BP was lowered below 70 mm Hg; no such nondiabetic patients (140/90 mm Hg) gave a target for diabetics pattern was seen for strokes. The consensus is that systolic BP is identical to that recommended in the earlier US hypertension more predictive for most patients, and the benefits of lowering guidelines: < 130/85 mm Hg! In the 1501 diabetics in the HOT systolic BP probably outweigh the risk of lowering diastolic BP study, the lowest risk of stroke, heart attack, or CVD death in the in most hypertensives, even with CHD. 100 intent-to-treat analysis was seen in those randomized to a diastolic BP < 80 mm Hg. 90 The most recent clinical trial data in favor of a lower BP target for diabetics comes from the Strong Heart Study, Different Targets for Blood Pressure Goals? which observed significantly lower CVD event rates among the diabetics ran domized to BP < 130/80 mm Hg or an LDLcholesterol target of < 100 mg/dL. 93

There have been concerns that the lower BP goal for diabetics would be expensive, as it requires more antihypertensive pills and more visits to health care providers, but several studies show this to be untrue. A cost analysis of UKPDS showed that despite higher drug and provider costs, the strategy of the lower BP goal saved lives, strokes, limbs, and money. The cost-effectiveness ratio for the lower BP goal was - £720 per year of life saved and an even more impressive - £1049 per year of life without diabetic complications.

JNC 7 recommended a BP target of < 130/80 mm Hg for patients with CKD, but there are few clinical trial data to support this. The most favorable long-term data came originally from the Modification of Diet in Renal Disease (MDRD) trial, which overrecruited patients with polycystic kidney disease but did show a benefit of BP < 125/75 mm Hg. 94 In a patient-level meta-analysis of placebo-controlled trials of ACE inhibitors in CKD, the optimal achieved systolic BP to avoid doubling of serum creatinine or endstage renal disease was 110 to 129 mm Hg. 95 In a direct comparison of BP targets for patients with hypertensive nephrosclerosis, the

American Study of Kidney Disease and Hypertension (AASK), **135** there was no significant benefit to reducing BP to < 125/75mm Hg (compared with < 140/90 mm Hg) during the first 3 years of follow-up. 96 Unfortunately, the 10-year incidence of doubling

antagonist, and the combination was used in the Controlled ONset of serum creatinine or end-stage renal disease among the cohort Endpoints initially randomized to the ACE inhibitor and the lower BP goal on the basis of data from MDRD, AASK, and observational studies. 3,85,95

In 2007, an American Heart Association Scientific State ment suggested that patients with established heart disease should be treated to < 130/80 mm Hg, and an "optional target" for those with heart failure is < 125/75 mm Hg. 40 These recommendations are controversial because few trials randomized subjects with heart disease to different target BPs and observed subsequent outcomes. to a lower than usual BP goal is analogous to the situation for diabetics 3-5,13,25,84 and those with CKD, 3-5,85 the clearest clinical Although the main trial was designed to compare outcomes in significant increase in atheroma volume, patients with BP For diabetic patients or those with CKD, JNC 7 recommends a between 120-139/80-89 mm Hg had no major change, and those

Recent research has suggested that traditional BP measures at the brachial artery correlate poorly with the BP experienced by the heart and brain, primarily because of disordered wave reflections, primarily in older patients with stiff arteries. ¹⁰² Many different noninvasive methods of measuring central aortic pressure have been developed, some of which are reimbursed by insurance. Several studies have shown a better prediction of CVD events with central aortic than with brachial BPs, 103 and most of the curvature in meta regression analyzes (see Fig. 9-5) disappears if estimates of central aortic systolic pressure are used on the *x* -axis rather than systolic BP. The largest clinical trial to prospectively measure central aortic pressure is the Conduit Artery Func tion Evaluation (CAFE), 69 a substudy of ASCOT. 68 A potential explanation for the impressive reduction in CVD events in ASCOT with the amlodipine-based regimen, despite a small difference in BP compared with the atenolol-based regimen, is that atenolol did not lower central aortic pressure nearly as well as amlodipine did. Sophisticated multivariate

analyzes demonstrated significant CVD benefits for every 10 mm Hg further lowering of central aortic pressure in CAFE. ⁶⁹ Whether technically challenging serial estimations of central aortic pressure should become a target for therapy or simply a risk stratification tool (particularly in older patients) is currently controversial. Proponents tout the pathophysiological links between the biophysics of aging artery ies, collagen crosslinking, the resulting low-capacitance vessels, ventricular-vascular coupling, and the atherosclerosis rhotic process as reasons to consider central aortic pressure an appropriate target for primary prevention. ^{102,104}

More sophisticated technologies have also resulted in non-invasive methods of measuring pulse wave velocity, -augmentation index, and aortic stiffness. ¹⁰⁵ Whereas their utility as 9 risk stratification tools has been demonstrated (most convincingly in the prospective Strong Heart Study), no clinical trials have yet used these measurements as targets for therapy. Because some of these parameters closely correlate with data available from 24-hour ABPM, ¹⁰⁶ it is likely that investigators will soon reexamine their ABPM files and begin finding more data to support these measures of vascular function as prognostic tools.

Constructing a Regime

Because most hypertensive patients will require two or more drugs to achieve the more intensive treatment goals outlined before, the health care provider probably should be concerned with choosing appropriate drug combinations more than with which specific agent is selected to begin the BP-lowering process. ^{5,9,15} Even in ALLHAT, which restricted enrollment to those with a high chance of being controlled on single-drug therapy (untreated BP < 180/110 mm Hg, or < 160/100 on one or two drugs), an average of two medications per patient was required at the end of the study. ⁸⁶

The ideal drug regimen would have many characteristics. In addition to lowering the BP to goal, the regimen should be able to be administered once daily without regard to meals, be relatively inexpensive, cause few adverse effects (and perhaps even result in fewer side effects than single-drug therapy), and be widely available in all pharmacies and ben efits plans. Several of the newer fixed-dose combination products have several of these attributes; those combining a dihydropyridine calcium antagonist and an ACE inhibitor, for instance, result in less pedal edema than the calcium channel blocker alone, even at the same doses. We now have three triple-drug combinations in the United States; one contains a diuretic, reserpine, and low-dose hydralazine; the other two combine a diuretic, amlodipine, and an ARB.

Management in Special Circumstances

Aside from patients who have a compelling indication for a particular antihypertensive drug (see Table 9-4), hypertensive patients having increased CVD risk require more attention. Those falling into this category include hypertensive patients with older age, diabetes, pregnancy, impending surgery, refractory hypertension, renal impairment, or hypertensive crisis.

Older Patients with Hypertension

There is now no question that older patients derive major benefit from antihypertensive drug therapy. ⁴¹ In fact, the number needed to treat (and similarly the cost) to prevent a single CVD event is *far* lower for older (compared with younger) patients, simply because their vintage puts them at higher absolute risk. Thus, they are also candidates for antiplatelet and lipid-lowering therapy, even if they do not have established CHD. ^{68,90} The decades-old controversy about whether hypertensive individuals older than 80 years benefit from antihypertensive drug therapy has been resolved in the affirmative by the HYpertension in the Very Elderly Trial

(HYVET). It was stopped after only 1.9 of the 5 years of planned follow-up because of a significant benefit on all cause mortality, despite a final nonsignificant benefit on stroke (the primary endpoint). ⁴²

Diabetic Patients with Hypertension

The regimen used to treat diabetics to a BP goal of < 130/80 mm Hg should include either an ACE inhibitor or an ARB (see Table 9-4). 3 5,13,14,25,36,84 An ARB can be justified if the patient is intolerant of an ACE inhibitor or if the patient has type 2 diabetes, renal impairment, and major proteinuria (eg, patients in the Irbesartan Diabetic Nephropathy Trial [IDNT] or the Reduction of Endpoints in Noninsulin dependent diabetes mellitus with the Angiotensin II Antagonist Losartan [RENAAL] trial). The Captopril Cooperative Study Group demonstrated renoprotective effects of an ACE inhibitor in type 1 diabetics. Either an ACE inhibitor or an ARB has reduced the progression of microalbuminuria to overt proteinuria (in the MIcroalbuminuria, Cardiovascular and Renal Outcomes substudy of the Heart Outcomes Prevention Evaluation [MICRO-HOPE] and the second IRbesartan MicroAlbu minuria [IRMA-2] trial). The cardiovascular benefits of an ACE inhibitor in type 2 diabetics have been seen in MICRO HOPE. The challenge for most diabetics is to achieve the several recommended goals (BP < 130/80 mm Hg, LDL-cholesterol < 100 mg/dL, A1c < 7%). Unfortunately, most surveys of large groups of patients have shown worse control of BP in diabetic subjects compared with nondiabetics. However, a randomized trial carried out at the Steno Diabetes Center in Denmark showed significant reductions in both CVD events and long-term mortality among those who were given the more intensive multifactorial intervention (which included BP < 130/80 mm Hg). 107

Pregnant Patients with Hypertension

Many of the antihypertensive drugs are either contraindicated or a potential threat to the mother or fetus. ¹⁰⁸ Diuretics, the "preferred" first-line therapy for nonpregnant hypertensives, are generally avoided during pregnancy owing to the risk of oligohydramnios. ACE inhibitors, ARBs, and renin inhibitors are contraindicated because of the risk of renal and other fetal malformations. Nitroprusside is transformed to cyanide, which is very toxic to the fetus. As a result, time-tested and traditional antihypertensive drug therapy is usually used, including, in order, methyldopa, hydralazine, a beta blocker (typically labetalol), and then perhaps a calcium antagonist.

Hypertensive Patients with Impending Surgery

Most hypertensive patients receive closer scrutiny when elective surgery is planned. On occasion, the procedure must be postponed, especially if the BP is > 160/95 mm Hg. Most anesthesiologists recommend holding the preoperative dose of an ACE inhibitor or ARB, as they increase the risk of hypo tension during induction. Adding a beta blocker to reduce perioperative CVD risk is not recommended unless it was given chronically before the procedure or the patient is having major vascular surgery. Even these recommendations have been criticized on the basis of a meta-analysis of 33 trials including 12,306 patients. ¹⁰⁹

Patients with Refractory Hypertension

Although several definitions of refractory hypertension have been proposed, the problem is so important that an American Heart Association Scientific Statement has recently summarized our knowledge on the topic. ⁵¹ The "cause" of resistant hypertension can usually be determined for a specific patient after a proper evaluation. The causes of resistance

TABLE 9-6	Estimated Prevalence of Common Causes of Resistant Hypertension in Primary Care Versus Referral Centers		
Primary Care	Cause of Resistant Hypertension	Tertiary Center	
~ 50%	Nonadherence to prescribed drugs	10%-20%	
20%-30%	Problems with blood pressure measurement White coat hypertension	t 10%-20%	
2070 0070	Medication-related causes	1070 2070	
~ 5% ~ 10% < 2%	Suboptimal medication regimen Drug- drug interaction Objective medication intolerance	30%-50% < 5% < 2%	
1%-5% 1%-7%	Interfering substances Excessive dietary sodium intake Ethanol, NSAIDs, or illicit substances	1%-10% 1%-12%	
< 5% < 5%	Secondary hypertension Sleep apnea, aldosterone excess Traditional secondary causes of hypertension	1%-83% 5%-28%	
1%-2% 1%-2%	Psychological causes Anxiety, panic disorder, depression Subjective medication intolerance	1%-2% 1%-5%	

NSAIDs, nonsteroidal anti-inflammatory drugs. Medication intolerance was judged to be "objective" if the patient-reported adverse effect had been previously noted in the literature or "subjective" if it had not been previously noted (eg, hair hurts, teeth itch).

Estimated from Calhoun 51 and other sources.

hypertension differs, depending on whether the patient is seen in a primary care setting or a referral center (Table 9-6). The most common successful pharmacological intervention for patients with resistant hypertension in many centers is modification of the drug regimen. Adding or switching to an appropriate diuretic (a loop diuretic if the eGFR is < 40 to 50 mL/ min/1.73 m 2 or thiazide-like diuretic otherwise) 52 and adding an aldosterone antagonist or an alpha blocker are the most frequent successful changes. Research has not generated very useful solutions for the all-too-common scenarios of the older, diabetic, or autonomously challenged patient with resistant systolic hypertension who develops orthostatic hypotension on treatment. Such patients sometimes respond to individualized treatment regimens (sometimes including venous compression stockings, low-dose midodrine, or dietary salt liberalization); more commonly, the BP target has to be modified upwards.

Hypertensive Patients with Chronic Kidney Disease

People with hypertension and eGFR < 60 mL/min/1.73 m ² (ie, stage 3 CKD ⁸⁵) should have their renally excreted anti hypertensive drugs reduced in dose (or, less commonly, frequency of administration) and be treated to a lower target (currently < 130/80 mm Hg). ^{3-5.85} An ACE inhibitor or ARB is highly recommended, after which a modest acute rise in serum creatinine (< 25% from baseline) and usually a clinically insignificant increase in serum potassium should be expected. An ARB delayed the progression of renal disease in type 2 diabetics in both IDNT and RENAAL ⁴; an ACE inhibitor was more effective than the calcium antagonist or beta blocker in AASK. ⁹⁶ An ACE inhibitor can be recommended for nondiabetic renal impairment. ^{86,95,110}

The role of albuminuria or proteinuria as a risk factor for CVD or as a target of therapy is fascinating and controversial. Many epidemiological studies (including normotensive and nondiabetic

subjects in the Framingham Heart Study 111) and a few clinical trials have demonstrated albuminuria to be a predictor of both progression of renal disease and incident CVD. 3.5,9,85,94.97,110 The US Food and Drug Administration (FDA) has never recognized albuminuria as a proper surrogate endpoint, despite the opinions of many physicians and nearly all nephrologists. As a result, no drugs are approved to reduce albuminuria, which is most commonly assessed today with an early morning urine sample sent for albumin/creatinine ratio. 85 Many clinical trials have nevertheless proven that BP lowering, ACE inhibitors, ARBs, and renin inhibitors are more likely than dihydropyridine calcium antagonists to reduce albuminuria. 112 The strongest data arguing that giving drugs to lower albuminuria reduces CVD risk may arise from the Losartan Intervention For Endpoint reduction (LIFE) trial,9 113 but teasing out which attributes of a particular drug regimen have the largest effect on specific CVD events is challenging.

Patients with Hypertensive Crises

Hypertensive emergencies are best treated in the hospital (typically in the intensive care unit) with a short-acting, rapidly reversible, intravenously administered antihypertensive drug (typically nitroprusside, although the short-acting dihydropyridine calcium antagonist clevidipine, which is hydrolyzed by serum esterases, has been recently approved). 114 Hypertensive emergencies may be routinely treated in the out patient setting with any one of a number of oral antihypertensive agents, including captopril, labetalol, and clonidine. Nifedipine capsules, which had been widely used in this setting for nearly 20 years, are now to be used "with great caution, if at all," according to an FDA advisory, because of their propensity to cause quick and excessive hypotension. Perhaps the most important aspect of the treatment of hypertensive urgency is to arrange quick follow-up for the patient where chronic, better management of BP can be ensured.

Organizing for Successful Management

The goal of hypertension management is to prevent the morbidity and mortality associated with it and to do so in the "least intrusive" manner (both physiologically and fiscally). Because hypertension is not a disease but a condition that increases CVD and renal risk, its long-term control is a continuing challenge. For many years, it was thought that most patients with hypertension have no noticeable symptoms. Recent studies using antihypertensive drugs without appreciable side effects have demonstrated a significant decrease in headache when hypertensive patients are successfully treated. The quality of life among treated hypertensives was greatest in the group that achieved the lowest BPs in both HOT 90 and the Treatment of Mild Hypertension Study, 81 suggesting that there may be subtle symptoms that can be attributed to an elevated BP that improve when BP is lowered. It is nevertheless often difficult to convince a person with hypertension that taking a pill or changing one's lifestyle will result in tangible benefits, especially in the short term. It is also unfortunately true that treating hypertension (even successfully) does not reduce CVD risk to the level of a normotensive person. This provides strong impetus for the initiation of lifestyle modifications early, even before the levels of BP we call hypertension are present.

It is difficult to motivate patients to sustain their lifestyle modifications and to adhere long term to their prescribed medications. National survey data indicate that only 50.1% of America's hypertensives have their BPs < 140/90 mm Hg ⁶; other countries have even lower control rates. Nonadherence

138 results in wasted health care resources (- 10% of costs for hypertension), unnecessary hospitalizations, preventable strokes and heart attacks, and a large portion of the admissions to nursing homes (where drug taking can be more fully and efficiently supervised). Many suggestions have been made to increase adherence to pill taking; some have been proven successful in clinical trials. Several clinical trials in conditions besides hypertension (eg, after myocardial infarction or in heart failure) have shown that patients who do not take the pills as directed typically suffer more I CVD events than those who are adherent. Education of the patient (and family) is the cornerstone of improving adherence; patients with educational or cognitive deficits about hypertension and its treatment are unlikely to follow instructions for very long. Some clinics have improved their hypertension control rates after a health educator was added to the hypertension treatment team. Behavioral suggestions are often useful, such as integrating pill taking into the activities of daily living (eg, taking pills when caring for teeth) or using a pill organizer (typically to organize pills according to days of the week on which they are to be consumed). Increasing social support appears to be a beneficial strategy, especially for older individuals. The family member or caretaker can remind the patient of the need to take pills and to keep office visits as well as actually measure BP with a home device.

Missing appointments for follow-up care and monitoring of hypertension treatment has been associated with poorer outcomes. Several routine procedures can help minimize this problem. Appointment reminders (either by telephone or by mail) increase return visit rates. Scheduling of a specific time and date with a known health care provider, at the end of the office visit, is more successful than "calling in for a future appointment." Decreasing waiting times, having convenient office hours, and having a solicitous, caring office staff are also helpful.

Several characteristics of physicians have an impact on patients' adherence to medications and willingness to keep appointments. Physicians who are willing to involve the patient (when appropriate) in medical decision making are more successful in controlling BP. A common example is asking whether the patient would prefer to take a less expensive pill more often or a more expensive pill just once a day. Physicians who are perceived as having effective communication skills, who encourage questions from the patient and appropriate family members, and who provide feedback about the patient's progress also achieve better results.

Much of the underachievement of BP control nationwide has been attributed to patients' unwillingness to take pills and to appear for follow-up visits, but some of the blame has been attributed to physicians and other components of the health care system. Most systems have accepted the treatment of hypertension as a worthwhile endeavor as it saves money in high-risk patients and is relatively cost-effective in others (compared with many other common medical interventions). Recent efforts to restrict pharmacy benefits, to limit the range and doses of drugs on an accepted formulary, and to reduce the accessibility of health care services have led to increased costs. System-wide efforts (often called disease management programs) to encourage acceptance of generic drugs, to increase the threshold for beginning antihypertensive drug therapy in low-risk patients, to use one drug to treat both hypertension and a concomitant medical condition, and to encourage adherence to medication taking have been more successful. The workload of the health care provider involved in these efforts can be increased by some of these procedures, but it is sometimes offset by case managers and other allied health professionals who perform some of these important tasks.

CONCLUSION

Hypertension control is an important public health goal that requires

a long-term commitment from the patient, the physician, and the health care system. When all work together towards a common goal, the benefits of hypertension therapy in clinical trials can be easily and effectively translated into practice. This will eventually reduce the burden of disease and adverse cardiovascular and renal outcomes that were formerly associated with untreated or undertreated hypertension.

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CHAPTER 10

Heart Failure Prevention

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KEY POINTS

- Heart failure prevalence continues to rise in the United States and is projected to become worse with aging of the population.
- Recent therapeutic advances in heart failure have translated into only modest improvement in outcomes at the community level.
- Prevention efforts for heart failure have significantly lagged behind research for new treatment options.
- Optimal risk stratification strategies for incident heart failure continue to evolve.
- Hypertension and coronary heart disease remain the most prominent risk factors for heart failure, although cardiometabolic risk factors (e.g., obesity and diabetes) are rapidly increasing in importance.
- Several individual risk factor treatments have been shown to reduce the risk for development of heart failure.
- Further research is needed to assess optimal multiple risk factor interventions, behavior modification strategies, and novel therapeutic approaches to reduce heart failure risk in individuals with risk factors.
- Population-based promotion of healthy lifestyle at the primordial prevention level is likely to have an impact on the largest at-risk population.

The concept of "prevention" encompasses a vast spectrum ranging from primordial to primary, secondary, and tertiary prevention. For heart failure, an example of primary prevention is avoidance of behaviors that enhance (eg, weight gain) and embracing of behaviors that mitigate (eg, active lifestyle) the risk for development of heart failure. Primary prevention includes control of heart failure risk factors (eg, hypertension). Secondary prevention efforts institute interventions to reduce the risk of worsening of existing heart failure (eg, treatment with beta blockers). Finally, tertiary prevention interventions in advanced heart failure are implemented to reduce the chances of imminent mortality (eg, implantation of mechanical circulatory assist devices).

Most heart failure quality improvement efforts (eg, the Get With The Guidelines program by the American Heart Association) targeted at currently secondary prevention. 1 Similarly, most of the research efforts in heart failure are also targeted at treatment. During the last two decades, an intensified focus on the prevention of coronary artery disease has included the development of risk prediction models, 2-4 assessment of novel biomarkers, 5,6 identification of subclinical markers of coronary risk, 7,8 design of therapeutic trials for coronary disease prevention, and establishment of specific guidelines. 9,10 Despite the worsening epidemic of heart failure, however, prevention efforts in this area remain rudimentary.

Given the current and projected societal toll of heart failure, both clinically and economically, a more concentrated effort on heart failure prevention is needed. In this chapter, we focus on the primordial and primary prevention of heart failure. In recognition of the emphasis of this text, we will not discuss interventions to improve outcomes in individuals with manifest heart failure.

HEART FAILURE EPIDEMIOLOGY

Incidence, Prevalence, Outcomes, and Costs

It is estimated that more than 5.5 million subjects in the United States have heart failure, and more than 650,000 are diagnosed for the first time each year. 11 Current evidence suggests that the two major clinical subsets of heart failure (ie, patients with impaired versus preserved left ven tricular systolic function) each compose about half of the overall burden of heart failure in the community. 12 However, various studies have suggested that the proportion of heart failure with preserved ejection fraction ranges from 13% to 74% of all heart failure. ¹³ These studies are -heterogeneous and differ considerably in patient selection criteria, diagnostic criteria for heart failure, and quantitative methods for assessment of ventricular systolic function. 14 A common finding among these studies, however, is that the proportion of heart failure with preserved ejection fraction increases with age. 12,15 Not surprisingly, there is a secular trend towards increased prevalence of heart failure with preserved ejection fraction among patients hospitalized for heart failure. 16

Heart failure is the primary reason for 12 to 15 million office visits and 6.5 million hospital days annually. Recurrent hospitalization is a major quality of life and cost issue; the annual number of hospitalizations now exceeds more than 1 million for heart failure as a primary diagnosis and 2.4 to 3.6 million as a primary or secondary diagnosis. Heart failure patients are particularly prone to rehospitalizations, and read mission rates are near 50% within 6 months of discharge. The number of heart failure cases and deaths attributable to heart failure has increased steadily despite advances in treatment and a decline in other major

cardiovascular disease during the same interval. Heart failure remains the most common Medicare diagnosis-related group, and more Medicare dollars are spent on heart failure than for any other diagnosis. It has been estimated that the total direct and indirect cost of heart failure in the United States exceeds \$30 billion. ¹¹

Despite this, the outcomes of these patients continue to remain suboptimal at the population level, with only approximately 50% of the individuals surviving past 5 years after diagnosis. ¹⁷ Quality of life remains poor. Some temporal improvement trends in outcomes have been restricted primarily to individuals with systolic dysfunction, with no major advances in therapy for either patients with heart failure and preserved ejection fraction or those who are hospitalized for decompensated heart failure.

10 Future Projections

Heart failure prevalence is rising, and this trend is expected to continue. This is attributed to the increasing elderly population , improved care of acute heart diseases, and increasing prevalence of several cardiovascular risk factors like diabetes and obesity. According to the White House Conference on Aging, aging of the 78 million baby boomers will result in 1 in 5 Americans older than 65 years by 2050. ¹⁸ This trend will have a significant effect on health, health care, and health care economics because the use of formal and informal services is strongly correlated with age.

Heart failure incidence and prevalence are highest among the elderly. The incidence rate approaches 10 per 1000 population - annually after the age of 65 years; 80% of patients hospitalized with heart failure are older than 65 years. Thus, the increasing age of the population is expected to significantly worsen the current heart failure epidemic. Much of heart failure research to date has focused on treatment. Considering the current epidemiological trends and the future projections, it is imperative to further heart failure prevention efforts aggressively.

RISK FACTORS FOR HEART FAILURE

Many studies have described various risk factors for the development of heart failure ranging from lifestyle factors to dietary habits, medications, laboratory and imaging characteristics, and novel biomarkers and genomic markers (Table 10-1). 19 Heart failure risk increases proportionally with advancing age. Male gender is associated with a higher risk and may in part be explained by the higher prevalence of coronary disease in men. ²⁰ Behavioral risk factors for heart failure include lower levels of physical activity, coffee consumption, and increased dietary salt intake. Low socioeconomic status has been associated with increased risk. 20 Older age, hypertension, diabetes, obesity, and coronary artery disease are risk factors for heart failure with either reduced or preserved ejection fraction. In general, patients with preserved ejection fraction are older, are more likely to be female, and are more likely to be obese. In heart failure with preserved ejection fraction, hypertension is a more common risk factor 16; however, a substantial proportion of patients with reduced ejection fraction also have a history of hypertension. ¹⁶ In heart failure with reduced ejection fraction, ischemic heart disease is the most common etiology, but it is also a comorbid condition among many patients with preserved ejection fraction. 16 Among patients admitted for decompensated heart failure, 63% of patients with reduced ejection fraction and 54% of patients with preserved ejection fraction have coronary artery disease. 21 Thus, from a prevention perspective,

TABLE 10—1 Risk Factors for Development of Heart Failure

Demographics Older age

Male gender

Socioeconomic status

Education

Comorbidities

Hypertension
Left ventricular hypertrophy

Prior myocardial infarction

besity

Diabetes mellitus

Valvular heart disease

Renal failure

Dyslipidemia

Sleep apnea

Tachycardia

Impaired lung function Depression, excessive stress

Pharmacological exposures

Chemotherapeutic agents

Nonsteroidal anti-inflammatory

Thiazolidinediones

D.

Doxazosin

Lifestyle factors

Cocaine

Tobacco use

Excessive alcohol consumption Excess caffeine

Excess sodium consumption

Echocardiographic parameters

Ventricular size Ventricular mass

Ventricular dysfunction Diastolic filling impairment

Biochemical markers

Albuminuria

Homocysteine
Tumor necrosis factor- a

Interleukin-6

C-reactive protein

Insulin-like growth factor 1 Natriuretic

peptides

Genetic risk factors

Adrenergic receptors a 2 c Del322-

325 deletion

Adrenergic receptors p 1 Arg389

change

Overexpression of protein kinase C- a ACE and Ang II type 1 receptor

gene polymorphisms

ACE, angiotensin-converting enzyme; Ang II, angiotensin II.

the major target risk factors for possible modification are common in both subsets of heart failure.

Hypertension and coronary artery disease are the most common and strongest risk factors, conferring a twofold to threefold increased risk for heart failure. Both elevated systolic blood pressure and elevated diastolic blood pressure have been associated with increased risk for heart failure. ²² Diabetes mellitus is associated with a higher risk, both with and without the presence of simultaneous coronary heart disease. Valvular heart disease increases the risk through hemo dynamic alterations on the ventricles, with either volume or pressure overload leading to myocardial dysfunction.

Obesity, through multiple mechanisms, predisposes individuals to heart failure. ²³ Excessive alcohol intake increases blood pressure and is a direct myocardial toxin ²⁴; however, light to moderate alcohol consumption has been inversely associated with heart failure risk, especially in men. ^{25,26} Smoking promotes several cardiovascular risk factors and is associated with heart failure development. ^{17,20} Dyslipidemia predisposes to heart failure; however, it is unclear if this is primarily related to atherosclerosis or if there are alternative effects. High total cholesterol, low high-density lipoprotein cholesterol, and high triglyceride levels have all been correlated with greater left ventricular mass and impaired diastolic function. ^{27,28} However, observational studies investigating the association of total cholesterol with incident heart failure in individuals without prior coronary heart disease have yielded inconsistent results. ^{29,30}

Complications of renal dysfunction, including anemia, hypertension, arterial stiffening, sodium and water retention, endothelial dysfunction, and alteration in biomarker profiles, are associated with elevation of the risk for development of heart failure. Anemia and sleep-disordered breathing are

heart failure odds; whether this represents a compensatory factors). The "independent" prognostic power of any risk factor response to lower stroke volume and underlying symptom atic depends largely on what it is compared against. Importantly, systolic dysfunction or neurohumoral activation, or both, is not despite strong etiologic association with a disease, a risk factor clearly known. Finally, several abnormalities of pulmonary - may be limited in its prognostic role. 35 function, including reduced forced vital capacity and forced expiratory volume in the first second of expiration, are associated disease than the usual etiological research if it is to provide a basis with heart failure risk.

myeloperoxidase levels 33 have been associated with increased risk sensitivity and specificity. 35 as well.

phosphamide, and 5-fluorouracil) are associated with heart failure. classification scheme that includes stage A patients, those who Doxorubicin-induced cardiotoxicity is dose related, especially do not have any structural heart disease but are at risk for heart when the cumulative dose exceeds 550 mg/m². Recent data show failure. ³⁶ This new classification is presented in Figure 10-1. a higher risk with the use of trastuzumab. Cyclooxygenase 2 Although individual risk factors for heart failure (eg, inhibitors may increase the risk of myocardial infarction, raising hypertension) are well described, how to quantify individual risk heart failure risk concerns. Thiazolidinediones have been in patients with various combinations of risk factors is not clearly associated with edema and precipitation of heart failure. Recently, described. the Rosiglitazone Evaluated for Cardiac Outcomes and Regulation of Glycaemia in Diabetes (RECORD) study showed that Framingham risk score, have been developed and validated for thiazolidinedione use is associated with a small but clinically coronary events. ² However, the heart failure syndrome

dilation with an increase in end-diastolic or end-systolic preceding coronary event before developing heart failure. 37 dimensions, increased left ventricular mass, left ventricular diastolic filling impairment, left atrial enlargement, asymptomatic systolic dysfunction.

Finally, there is growing interest in discovering the genomic predictors of heart failure. 19 Genetic alterations in functional Most studies of heart failure risk factors have targeted individual pathways, such as energy production and regulation (eg, risk factors. Table 10-2 summarizes the nine studies that have mitochondrial mutations), calcium cycling abnormalities (eg, RYR2 assessed independent risk factors for incident heart failure mutations), and mutations in transcriptional regulators (eg, Nkx2.5 comparing multiple risk factors 20,37-44; only two of them developed leading to ventricular hypertrophy), may lead to heart failure risk. a prediction model for incident heart failure. 40,44 The Framingham Genetic polymorphisms in sympathetic receptors, such as the genes heart failure risk score assessed the probability of developing heart coding for alpha _{2C} -adrenergic receptors (a _{2C} Del322-325) or beta failure during a 38-year period in which there were 6354 person-1 -adrenergic receptors (B 1 Arg389), are associated with heart examinations in men and 8913 in women. 40 Although regression failure risk. Interestingly, homozygous blacks for a 2C Del322-325 coefficients differed between men and women somewhat, overall have a fivefold higher risk. If this polymorphism is associated with predictors were similar (see Table 10-2). Interestingly, systolic homozygosity for \$ 1 Arg389, the risk increases by 10-fold. Other blood pressure was predictive only in men and diabetes mellitus in polymorphisms associated with heart failure risk factors (eg, hypertension and vascular stiffness) have angiotensin-converting enzyme and AT₁ receptor gene polymorphisms, which in turn may affect heart failure risk.

PREDICTING INCIDENT FAILURE Identification of High-Risk Individuals

For implementation of cost-effective preventive interventions, heart failure risk factors have been described, determination of other cohorts, such as the Health, Aging, and their role in predicting a future event is nevertheless challenging. A variable associated with a disease may be a risk marker or a risk factor. A risk factor's relative importance may change over time as the population

characteristics change (eg, secular trends in age, treatment 143

associated with heart failure. An increased heart rate increases modalities of other risk factors, and prevalence of competing risk

A risk factor must have a much stronger association with for detection and prediction in an individual patient. If the Microalbuminuria is associated with a threefold increased heart distributions of the risk factor differ between two groups of patients failure hospitalization risk. Levels of homocysteine, insulin-like with differing outcomes, the risk factor may be • statistically growth factor, proinflammatory cytokines (eg, tumor necrosis associated with the outcome. However, for the risk factor to factor- a , interleukin-6, and C-reactive protein), and B-type perform well as a prognostic test, the distributions in the two natriuretic peptide are associated with an increased risk. Recently, groups must be sufficiently separated to permit selection of a cutoff serum resistin, 31 lipoprotein-associated phospholipase A 2, 32 and value that will discriminate - Q between the two groups with high

The American College of Cardiology and the American Heart Several chemotherapeutic agents (eg, doxorubicin, cyclo - Association (ACC/AHA) recently proposed a new heart failure

Multiple risk factor prediction schemes, such as the relevant increased risk of heart failure. 34 However, the debate represents a spectrum ranging from ischemic to nonischemic continues as to whether these drugs truly predispose individuals causes and normal to depressed ejection fraction. Heart failure may to the development of heart failure or only promote fluid retention develop in elderly subjects as a result of age-related cardiovascular in individuals with prevalent left ventricular dysfunction. changes in the absence of traditional risk factors. High-risk subjects Recreational drugs (eg, cocaine abuse) may precipitate heart therefore may not be detected by coronary risk schemes. For example, in the Cardiovascular Health Study, 66% of incident heart Several cardiac anatomical and physiological measures are failure cases developed in subjects without baseline history of associated with a higher risk, including ventricular chamber coronary heart disease, and more than half of them never had a

Heart Failure Risk Prediction Models

women.

The investigators subsequently developed a 4-year event score with an event rate averaging 3.97 per 100 person-year examinations in men and 2.63 in women with a 37% increase per decade of age. There are several limitations to the use of this score. The chest radiograph and pulmonary function test requirement makes it more difficult to implement it in large population settings for screening. The model was drawn primarily on white populations, and its performance in other racial groups is not known. The model was derived on a restricted group of subjects with a known history of hypertension, coronary heart disease, or valvular heart disease; identification of high-risk individuals is essential. Although many such characteristics represented less than 50% of the population in

Stage A
At high risk for heart
failure but without
structural heart disease
or symptoms of heart
failure

Patients with: u

Atherosclerotic disease

Hypertension

Obesity

Diabetes mellitus

Metabolic syndrome

Using cardiotoxins

cardiomyopathy

With family history of

Patients

Structural heart disease

eart >

disease

Stage C
Structural heart disease with previous or current symptoms of heart failure

Stage D
Refractory heart failure requiring special interventions

Patients with: symptoms
Previous myocardial infarction

Left ventricular remodeling (left ventricular hypertrophy/low ejection fraction) Asymptomatic valvular

Stage B

Structural heart disease

but without symptoms

or signs of heart failure

heart failure symptoms

Patients with:

Known structural heart

Refractory

disease
AND
Dyspnea, fatigue,
reduced exercise
tolerance

Patients: Who has marked symptoms at rest despite maximal medical therapy

FIGURE 10-1 American Heart Association and American College of Cardiology new heart failure classification.

Development

of heart

failure

References	Cohort	Risk Factors
Eriksson, et al ∞	Gothenburg, Sweden	Hypertension, smoking, weight, heart size, T wave abnormality, heart rate variability, peak expiratory flow rate, and psychological stress
Chen, et al 39	EPES	Gender, age, diabetes, pulse pressure, and body mass index
Kannel, et al 40		
	Framingham Heart Study	Age, blood pressure, LVH, vital capacity, heart rate, CHD, murmurs, diabetes, cardiomegaly, and body mass index
Gottdiener, et al 3	Cardiovascular Health Study	У
		Age, gender, cerebrovascular disease, diabetes, blood pressure, FEV 1, creatinine, C-reactive protein, ankle-arm index, atrial fibrillation, LVH, abnormal ejection fraction, electrocardiographic ST-T abnormality
He, et al 20	NHANES I	Gender, education, physical activity, smoking, weight, hypertension, diabetes, valvular disease, and CHD
Wilhelmsen, et al.	41 Gothenburg, Sweden	Age, family history of infarction, diabetes, history of chest pain, smoking, coffee consumption, alcohol abuse, blood pressure, and body mass index
Bibbins-Domingo,	, et al. • Heart and Estrogen/Progestin	Diabetes, atrial fibrillation, myocardial infarction, creatinine clearance, blood pressure, smoking, body mass index, left bundle branch block, and LVH
Carr, et al 43	Replacement Study	
oan, or ar	RENAAL and LIFE studies	Age, history of myocardial infarction, vascular disease, atrial fibrillation, urinary albumin/creatinine ratio, alcohol abuse Cornell product, and body mass index
Butler, et al 4	Health ABC	Age, CHD, smoking, blood pressure, heart rate, serum creatinine, fasting glucose, albumin level, and LVH on electrocardiogram

CHD, coronary heart disease; EPESE, Established Populations • for Epidemiological Studies of the Elderly, FEV |, forced expiratory volume in the first second; LIFE, Losartan left Intervention For Endpoint reduction in hypertension; LVH, ventricular hypertrophy; NHANES, National Health and Nutrition Examination Survey; RENAAL, Reduction of the Endpoints in Non-insulin dependent diabetes mellitus with Angiotensin II Antagonist Losartan.

Body Composition (Health ABC) Study, ⁴⁴ limiting the model's generalizability. Finally, this model was not externally validated in independent cohorts.

Recently, the Health ABC heart failure risk model was developed with the data from 2935 individuals participating in the Health ABC Study. The mean age of the population was 73.6 years, with 52% females and 41% blacks. Independent predictors of heart failure included age, history of coronary heart disease and smoking, systolic blood pressure and heart rate, serum glucose concentration, creatinine and albumin levels, and electrocardiographic left ventricular hypertrophy; the model has good discrimination (C-statistic 0.73 in the derivation data set and 0.72 by internal validation with optimism-correction) and

good calibration. A simple point score was created to predict incident heart failure risk in four risk groups corresponding to < 5%, 5% to 10%, 10% to 20%, and > 20% 5-year risk (Fig. 10-2).

The investigators subsequently externally validated the model in the Cardiovascular Health Study; the model retained adequate predictive capabilities. ⁴⁵ The model predicted risk equally well in both men and women and in white and black races (Fig. 10-3). The utility of the Framingham heart failure risk score in predicting incident heart failure in the overall Health ABC cohort and the subcohorts in which the original score were developed (ie, subjects with known baseline history of hypertension, coronary heart disease, or valvular

Age		
Years	Points	
< 71	-1	
72-75	0	
76-78	1	
> 79	2	

Coronary Art	ery Disease
Status	Points
No	0
Possible	2
Definite	5

1
its

Systolic Bloc	od Pressur
mm Hg	Points
< 90	-4
95-100	-3
105-115	-2
120-125	-1
130-140	0
145-150	1
155-165	2
170-175	3
180-190	4
195-200	5
> 200	6

Heart Rate		
Bpm	Points	
< 50	-2	
55-60	-1	
65-70	0	
75-80	1	
85-90	2	
> 95	3	

Smoking			
Status	Points		
Never	0		
Past	1		
Current	4		

Alb	umin
g/dL	Points
> 4.8	-3
4.5-4.7	-2
4.2-4.4	-1
3.9-4.1	0
3.6-3.8	1
3.3-3.5	2
< 3.2	3

Health ABC HF Risk Score	HF Risk Group	5-yr HF Risk
< 2 Points	Low	< 5%
3-5 Points	Average	5-10%
6-9 Points	High	10-20%
> 10 Points	Very high	> 20%

 Mg/dL
 Points

 < 80</td>
 -1

 85-125
 0

 130-170
 1

 175-220
 2

 225-265
 3

 > 270
 5

Creat	inine
mg/dL	Points
< 0.7	-2
0.8-0.9	-1
1.0-1.1	0
1.2-1.4	1
1.5-1.8	2
1.9-2.3	3
> 2.3	6

Key:
Systolic BP to nearest 5 mm Hg
Heart Rate to nearest 5 bpm
Albumin to nearest 0.1 g/dL
Glucose to nearest 5 mg/dL
Creatinine to nearest 0.1 mg/dL

FIGURE 10-2 The Health ABC Heart Failure (HF) Risk Score. (From Butler J, Kalogeropoulos A, Georgiopoulou V, et al: Incident heart failure prediction in the elderly: the Health ABC Heart Failure Score. Circ Heart Fail 1:125, 2008. Reprinted with permission of Wolters Kluwer Health.)

heart disease) was also assessed. The Framingham heart failure risk score performance was inferior compared with the Health ABC heart failure risk model.

Challenges in Predicting Incident Heart Failure

Several unique issues make heart failure risk assessment challenging. Heart failure is a clinical diagnosis and cannot be easily diagnosed with a "test." This leads to diversity in opinions and diagnostic uncertainty in a certain proportion of cases. The most common clinical criteria used to diagnose heart failure are the Framingham criteria, which require the presence of at least two major criteria or one major criterion and two minor criteria. 46 Major criteria include paroxysmal nocturnal dyspnea, neck vein distention, rales, radiographic cardiomegaly, acute pulmonary edema, S₃ gallop, increased central venous pressure > 16 cm H₂O, circulation time > 25 seconds, hepatojugular reflux, or pulmonary edema or vis ceral congestion or cardiomegaly at autopsy. Minor criteria include bilateral ankle edema, nocturnal cough, dyspnea on ordinary exertion, hepatomegaly, pleural effusion, reduced vital capacity by one third from maximum, and heart rate > 120 beats/min. Investigators from the Cardiovascular Health Study developed alternative criteria that included medication use and imaging modalities. ⁴⁷ When both sets of criteria were compared, only half the patients were adjudicated to have heart failure by both criteria, whereas the other half were labeled with either one or the other but not both criteria. 48 Similar discordance has also been shown between administrative discharge diagnoses and designation based on detailed chart review. 49

Part of this discordance is related to the diagnosis of heart

failure with preserved ejection fraction. Many of the heart failure symptoms (eg, shortness of breath) and signs (eg, edema) are nonspecific and can be seen in other conditions (eg, obesity and chronic lung disease). The European Society of Cardiology developed a consensus statement for diagnosis of this condition by use of biomarker- and imaging-based detailed protocols. ⁵⁰ Similarly, another set of detailed clinical criteria to diagnose incident heart failure in clinical trials was recently published. ⁵¹ These criteria may be of limited usefulness from a population health perspective because of cost and logistics. Thus, these guidelines are likely to be used primarily in the clinical and research setting and not in the screening and population prevention arena.

Any clinical prediction rule is unlikely to diagnose "niche" heart failure, such as amyloidosis and hypertrophic cardiomyopathy, disorders with a distinct natural history. Simi larly, whether risk prediction for heart failure differs for low versus preserved ejection fraction or stage A versus stage B heart failure is not known.

Risk Factors and Population Attributable Risk

Assessment of the population attributable risk for the common risk factors for any disease is key to prioritize prevention

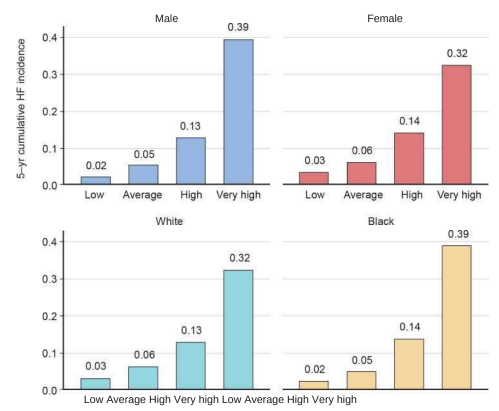


FIGURE 10-3 Health ABC Heart Failure Risk Score performance in gender- and race-stratified subgroups. Numbers represent fraction of participants with incident heart failure (HF). (From Butler J, Kalogeropoulos A, Georgiopoulou V et al: Incident heart failure prediction in the elderly: the Health ABC Heart Failure Score. Circ Heart Fail 1:125, 2008. Reprinted with permission of Wolters Kluwer Health.)

efforts cost-effectively. Population attributable risk represents the proportional reduction in disease risk that would be achieved by eliminating the risk factor from the population, assuming a causal relationship. The relative importance of risk factors in the population can help plan public health interventions; however, the absolute estimates for the population attributable risk of the various risk factors are highly dependent on the definition of risk factors and inclusion of other risk factors in the model used to derive the population attributable risks.

In a recent report from the Health ABC Study, ² a cohort of well-functioning, community-dwelling older adults, coronary heart disease and uncontrolled blood pressure were the leading causes of heart failure in whites and blacks and in men and women. A substantial proportion of heart failure, however, was also attributed to metabolic and cardiorenal factors, including glucose and renal abnormalities. Several previous investigations have reported substantial sex- and race-related differences in population attributable risks, disease development and progression, and prognosis for heart failure. ^{17,52} Understanding and quantifying these differences are important for planning appropriate preventive interventions.

The higher incidence of heart failure in black compared with white participants in the Health ABC Study was simultaneously accompanied by a higher prevalence of risk factors in black participants (Table 10-3). Interestingly, the black participants had a higher proportion not only of overall risk factors but also specifically of those risk factors that are potentially amenable to intervention, which translated the population attributable risk in general to a higher modifiable fraction in black individuals (68% versus 49% in whites). These data provide valuable information into race-based differences and help prioritize interventions, set targets, and assess the feasibility of novel therapies.

HEART FAILURE INCIDENT RISK MODULATION

Incident heart failure risk modulation is currently targeted primarily towards risk factor management individually. For coronary artery disease, the patient's treatment plan is tar geted on the basis of the individualized risk profile related to the various combinations of risk factors (eg, blood pressure goals or low-density lipoprotein level). On the contrary, no such paradigm for heart failure risk modulation currently exists. Many heart failure risk factors (eg, age and gender) cannot be intervened on. Other risk factors like proinflammatory states may be targets for intervention in the future. This section describes currently known interventions that directly or indirectly reduce the risk of incident heart failure. Most of these also affect other adverse cardiovascular outcomes; however, this section is focused primarily on heart failure-related data.

Lifestyle

Several studies have reported a reduced risk of incident heart failure with a healthy lifestyle. Maintaining a healthy weight, avoiding smoking, engaging in regular exercise, and maintaining a healthy diet have been shown to favorably influence heart failure risk factors, including coronary heart disease, ^{53,54} diabetes mellitus, ⁵⁵ and hypertension. ⁵⁶ Recently, the Physicians' Health Study investigators reported that healthy life style habits (ie, normal body weight, not smoking, regular

TABLE 10—3 Multivariable Rate Ratios and Population Attributable Risks for Clinical Risk Factors of Incident Heart Failure in the Health ABC Study

	White (n =	1686)	Black (n =	1167)
Risk Factors	RR (95% CI)	PAR (%)	RR (95% CI)	PAR (%)
Modifiable				
Systolic blood pressure >140 mm Hg	1.80 (1.27-2.55)	21.3	1.95 (1.33-2.84)	30.1
Coronary heart disease	2.72 (1.89-3.90)	23.9	3.31 (2.26-4.85)	29.5
Glucose >126 mg/dL	2.08 (1.35-3.22)	11.3	1.37 (0.88-2.14)	7.3
Left ventricular hypertrophy	0.90 (0.44-1.84)	_	2.20 (1.47-3.30)	19.5
Current smoking	2.04 (1.15-3.64)	5.5	2.08 (1.37-3.16)	15.0
Modifiable Fraction		48.9		67.8
Potentially Modifiable eGFR <60 mL/min/1.73m ²				
	1.29 (0.88-1.87)	6.8	2.14 (1.42-3.24)	16.2
Albumin <3.8 g/dL	1.46 (0.98-2.16)	8.5	1.63 (1.09-2.44)	12.7
Heart rate >75 beats/min	1.45 (0.94-2.23)	6.7	1.97 (1.30-2.99)	15.7
Potentially Modifiable Fraction		20.5		38.6

eGFR, estimated glomerular filtration rate; PAR, population attributable risk.

Note: Population attributable risks are not additive and do not add up to 100%.

From Kalogeropoulos A, Georgiopoulou V, Kritchevsky SB, et al: Epidemiology of incident heart failure in a contemporary elderly cohort: the health, aging, and body composition study. Arch Intern Med 169:708, 2009. Reproduced with permission of the American Medical Association.

exercise, moderate alcohol intake, consumption of breakfast cereals, and consumption of fruits and vegetables) were associated with a lower lifetime risk of heart failure, with the highest risk of 21.3% in men adhering to none of these life style habits and the lowest risk of 10.1% in men adhering to four or more of these habits. 57

Overweight and Obesity

Body mass index is associated with heart failure in a positive 23.58 and linear 23 fashion in both sexes. Although body mass index in the obese range (> 30 kg/m²) is clearly associated with an increased risk for heart failure, 23.58 there is controversy about body mass index in the overweight range (25 to 29.9 kg/m ²). ⁵⁸ Recent data, however, support that excess weight is also associated with heart failure. 57,59 Abdominal obesity may be a stronger predictor for heart failure than total obesity, 60,61 even in the absence of coronary heart disease. 62 In population studies, a strong association has been found between abdominal adiposity and features of metabolic syndrome, insulin resistance, and inflammation, 63-65 all of which have been related to heart failure. Individuals with abdominal obesity have more elevated sympathetic neural activation, 66 and visceral adipose tissue has higher expression of angiotensinogen than subcutaneous adipose tissue does. 67

The anatomical location of excess fat in the abdominal cavity may also be important as there is emerging evidence that increased intra-abdominal pressure leads to cardiac abnormalities that predispose to heart failure. 68 On the contrary, however, there are also data supporting no predictive differences between total and abdominal obesity (as evaluated by waist circumference), 69 and the association between obesity (total or abdominal) and incident heart failure may be mediated partly by insulin resistance. ⁶⁹ In addition, the relationship between obesity and risk of heart failure appears to become less important with age. 70 This age-related attenuation of the obesity and heart failure association needs further investigation.

Several mechanisms by which elevated body mass index increases the risk of heart failure have been proposed. 71-74 Figure 10-4 summarizes these mechanisms.

Alterations in cardiac loading. Obesity is associated with hemodynamic overload with increased blood volume and

- cardiac output. In addition, left ventricular afterload is elevated because of both increased peripheral resistance and greater arterial stiffness.
- Changes in cardiac structure and function. Both over weight and obesity are associated with increased left ven tricular mass, wall thickness, and dimensions, whereas longer duration and severity of increased weight accelerate remodeling. Obese individuals may fail to increase their ejection fraction with exercise secondary to abnormal diastolic function. Right ventricular afterload may be increased because of sleepdisordered breathing and left ventricular changes. Obesity is also associated with left atrial enlargement 75 and atrial fibrillation. 76 Hypertension is common in obesity and further worsens heart failure risk. In experimental studies, obesity with myocardial steatosis lipotoxicity lipoapoptosis, which have been linked to cardiac dilation, reduced contractility, and diastolic dysfunction. Cardiac metabolic changes include reduced glucose and increased fatty acid oxidation; these changes increase myocardial oxygen consumption and decrease cardiac efficiency. 74
- Activation of neurohumoral and inflammatory pathways. Obesity is associated with neurohumoral activation, renal sodium retention, and increased systemic and myocardial oxidative stress. 71,74,77 Sympathetic and renin-angiotensin activation directly through adipose tissue signals is common in obesity. Adipose tissue is a source of pro-inflammatory cytokines, such as tumor necrosis factor -a , interleukin-6, and C-reactive protein; these cytokines, which suppress cardiac function, have been associated with incident heart failure. 78,79
- Promotion of atherogenic conditions. Obesity is associated with hypertension, insulin resistance, diabetes mellitus, and dyslipidemia, all of which enhance the risk of myocardial infarction 80 and also mediate or directly cause heart failure.
- Predisposition to sleep-disordered breathing. Along with right ventricular changes, obstructive sleep apnea could occur

also lead to left ventricular hypertrophy related to and volume and filling pressures, and improves diastolic

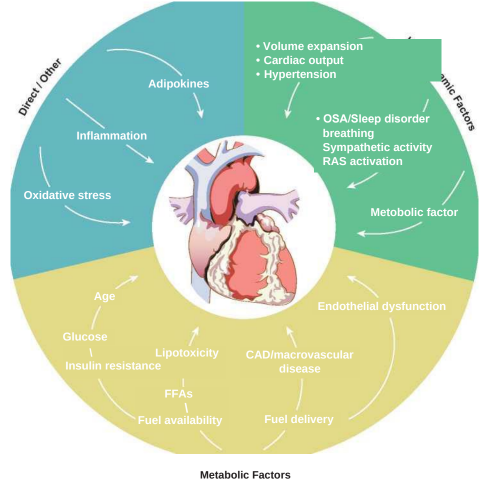


FIGURE 10-4 Multifactorial relationship between obesity and heart failure risk. CAD, coronary artery disease; FFAs, free fatty acids; OSA, obstructive sleep apnea; RAS, renin-angiotensin system. (From Abel ED, Litwin SE, Sweeney G: Cardiac remodeling in obesity. Physiol Rev 88:389, 2008. Reprinted with permission of the American Physiological Society.)

exacerbation of hypertension, increased sympathetic tone, chronic hypoxemia, and exaggerated intrathoracic pressure during obstructive episodes. 81

• Chronic kidney disease. Obesity is associated with an increased risk of proteinuria and renal insufficiency, presumably caused by glomerular hyperfiltration, glomerular hyperfusion, glomerular hypertrophy, hyperlipidemia, and increased expression of vasoactive and fibrogenic substances (such as angiotensin II, insulin, leptin, and transforming growth factor-S 1), all factors associated with heart failure risk. ⁷¹

Interventions. The main approach to cardiovascular risk reduction in obese patients should include weight control, physical activity, and control of the associated risk factors, such as hypertension, diabetes mellitus, sleep disorders, and components of the metabolic syndrome. ⁷³ Myocar dial changes with nonsurgical or surgical weight loss are feasible, and minor weight loss is efficacious; a 10% weight reduction improves systolic dysfunction, and weight loss of 8 to 10 kg produces a significant decrease in left ventricular dimensions and mass index and improves diastolic function.

Substantial weight loss reduces left ventricular wall thickness

measures and left ventricular systolic function. The hemodynamic benefits of weight reduction are important and further improve ventricular structure and function related to improved ventricular loading conditions. ⁷³The role of metabolic and neurohumoral modification may take precedence over the hemodynamic effects as left ventricular mass or functional improvement occurs independently of loading alterations. ⁸² Although many metabolic and neurohumoral interventions have been implicated in animal models, the roles of the reninangiotensin-aldosterone system antagonists, lipid-lowering therapy, and insulin-sensitizing drugs in obese humans need further study.

Sedentary Exercise Habits

Physical inactivity is an important risk factor for cardiovascular diseases including heart failure. ⁵⁹ Evidence suggests that regular physical activity has important and wide-ranging health benefits like reduction in risk of cardiovascular dis- eases, ⁸³ hypertension, ⁸⁴ and diabetes. ⁸⁵ Sitting more and per forming less nonexercise activity push the risk curve upward to the left as shown in Figure 10-5, where there is the most



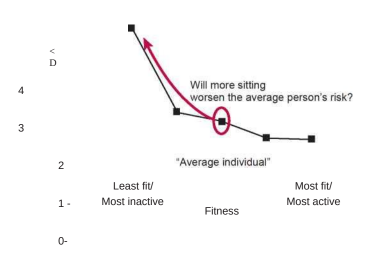


FIGURE 10-5 Sedentary lifestyle and risk for mortality. (From Hamilton MT, Hamilton DG, Zderic TW: Role of low energy expenditure and sitting in obesity, metabolic syndrome, type 2 diabetes, and cardiovascular disease. Diabetes 56:2655, 2007.)

risk for disease. Interestingly, emerging evidence indicates that maintaining a high level of daily low-intensity activity may be important independently of moderate to vigorous physical activity for several risk factors for coronary heart disease, such as elevated glucose, type 2 diabetes mellitus, and lipids such as triglyceride and high-density lipoprotein cholesterol levels, 86 all of which predispose to new-onset heart failure. Studies have linked prolonged sitting with car diovascular risk independently of age or recreational energy expenditure. 83.87

Physical activity is a key determinant of good health and an important component of weight reduction and weight maintenance, 88,89 improved lipoprotein profile, 88,90,91 and reduced risk of hypertension, ^{88,91,92} diabetes mellitus, ⁵⁵ and coronary artery disease. 88,93 These favorable influences on cardiovascular risk profile in turn reduce the likelihood of heart failure.

Physical activity could also reduce left ventricular hypertrophy independent of body weight or blood pressure because of reduction in vascular resistance and blood volume, improved endothelial function, suppression of the renin angiotensin and sympathetic nervous system activity, and reduction of insulin resistance. 91 Chronic physical activity reduces cytokine production by adipose tissue, skeletal muscles, endothelial cells, and blood mononuclear cells and upregulates antioxidant enzymes. 94 These modifying 104 to improve insulin sensitivity, 103 to lower plasma levels of effects on heart failure risk factors or intermediate pathways leading to heart failure can reduce incident heart failure.

Interventions. The integration of physical activity into the daily lives of the population has proven challenging. In the United States, from 2001 to 2005, the prevalence of regular physical activity increased by 8.6% (from 43.0% to 46.7%) among women and by 3.5% (from 48.0% to 49.7%) among men. 95 The recommendations of the American College of Sports Medicine and the American Heart Association for regular physical activity in healthy adults from 18 to improved endothelial function. 65 years currently include the following. 89

- · Aerobic activity. Moderate-intensity aerobic physical activity for a components of beverages on the risk of heart failure, whereas activity can be performed to meet this recommendation.
- · Muscle strengthening activity. It is recommended that 8 to 10

exercises be performed on two or more nonconsecutive days each49 week using the major muscle groups. To maximize strength development, a resistance (weight) should be used that allows 8 to 12 repetitions of each exercise resulting in voluntary fatigue. Muscle strengthening activities include a progressive weighttraining program, weight bearing calisthenics, stair climbing, and similar resistance exercises that use the major muscle groups.

· Activity dose. Vigorous-intensity activities may have greater benefit than moderate-intensity physical activity.

Because walking is the preferred activity among sedentary individuals embracing physical activity and the effects of walking have been reported as beneficial in primary prevention, % this should be adopted by individuals who do not adhere to the current 10 recommendations.

Alcohol Consumption

Excessive alcohol consumption regardless of beverage type is associated with alcoholic cardiomyopathy. 24,97 This entity is characterized by left ventricular dilation, increased mass, and reduced or normal wall thickness among patients with a lasting history of heavy alcohol consumption. ⁹⁷ Limited data are available on the amount and duration of consumption; most studies report that patients with symptomatic heart failure had 10 years or more of exposure to heavy drinking. 97

In animal models, excessive alcohol consumption is associated with left ventricular myocyte loss, negative inotropic effects and dysfunction of myocytes through abnormalities in calcium homeostasis, and hypertrophy through activation of cardiac betaadrenoceptors from the elevated levels of norepi nephrine. In humans, acute ethanol ingestion may also lead to depressed contractility. However, besides the direct myocardial toxicity, excessive alcohol consumption increases the risk of heart failure by promoting hypertension, myocardial infarction, 54 and diabetes. 98

Interestingly, other data are consistent with possible beneficial effects of moderate alcohol consumption on the risk of heart failure. The New Haven Epidemiologic Study of the Elderly program and the Cardiovascular Health Study reported a 47% ²⁵ and a 34% ⁹⁹ lower heart failure risk, respectively. The Framingham Heart Study reported a 59% lower risk among men who consumed 8 to 14 drinks per week compared with abstainers and only a modest and insignificant association in women. 100 Moreover, it has been reported that light to moderate alcohol consumption is associated with 40% to 50% lower risk of heart failure with previous myocardial infarction, whereas the risk of heart failure without antecedent myocardial infarction among heavy drinkers was 1.7fold higher than in abstainers in the same study. 101

Similar findings were reported in the Physicians' Health Study. ²⁶ Beneficial effects of alcohol have also been reported on risk for hypertension, 102 myocardial infarction, and diabetes mellitus, 103 whereas alcohol seems to raise high-density lipoprotein cholesterol, inflammatory markers and coagulation factors, and to raise plasma levels of adiponectin. 24 Furthermore, alcohol consumption has diuretic effects, which could prevent volume overload and delay the onset of signs and symptoms of heart failure. Beneficial effects of light to moderate alcohol consumption on incidence of heart failure seem to be independent of the reduced incidence of myocardial infarction and could be linked to lower pulmonary artery wedge pressure, reduced afterload, systemic arterial vasodilation, and

Current evidence does not support a major role for noneth anol minimum of 30 minutes on 5 days each week or vigorous- drinking patterns play an important role. Binge drinking (defined intensity aerobic activity for a minimum of 20 minutes on 3 days as consumption of three or more alcoholic drinks within 1 to 2 each week. A combination of moderate- and vigorous-intensity hours) has deleterious health effects, whereas light to moderate alcohol consumption (one or two

150 drinks per day for men and one drink per day for women spread over several days of the week) appears to yield most of the beneficial health effects. Thus, for a given volume of alcohol within the moderate drinking range, it is better to be distributed evenly throughout the week than to be consumed more rapidly.

Dietary Habits

In the Dietary Approaches to Stop Hypertension (DASH) diet, individuals are encouraged to consume more (1) fruits and I vegetables, (2) grains and grain products, (3) lean meats, fish, and poultry, (4) low-fat or nonfat dairy foods, and (5) nuts, seeds, and legumes and to reduce the consumption of red meat, fat, and sugar while maintaining a low-sodium intake.

recent evidence supports its beneficial effects on the reduction rate in women who adhere to the DASH diet. 105

The DASH diet may contribute to heart failure prevention in some cases because of reduction in blood pressure 56,106 and incident coronary heart disease. 56,107,108 Particularly women with the highest values of a score designed to measure consistency with DASH had a 24% lower risk of coronary heart disease and an 18% lower risk of stroke 107; using a different DASH score, Folsom and colleagues 108 did not find DASH had an 18% lower rate of death from coronary heart disease and a 14% lower rate of death from stroke that is similar to the projected effect from the DASH trial. 56,108 cholesterol levels and oxidative stress and exerts additional beneficial physiological effects like estrogenic effects of phytochemicals. 105

The relationship between several components of the DASH individuals, daily consumption of whole-grain breakfast cereals was associated with a 30% lower rate of heart failure compared with no consumption, 109 consumption of eggs more than twice per day was associated with a 64% higher rate, ¹¹⁰ consumption of fish was associated with a 20% to 31% lower heart failure rate depending on the frequency of consumption, 111 and consumption of 100 mmol or more of sodium was associated with a 26% higher rate 112; only nut consumption 113 was not associated with an increase or decrease in heart failure. In rat models of heart failure, macronutrient intake modified the course of cardiac dysfunction. High-fat diets reduced cardiac remodeling and contractile dysfunction; however, animals fed diets high in linoleic acid survived longer than those fed diets high in carbohydrates or lard. 114,115 When high-starch, high-fructose, worse survival. 116

There are several mechanisms by which whole-grain cereals can protect against heart failure risk, partially through effects on weight, hypertension, myocardial infarction, and diabetes mellitus. Nutrients contained in whole-grain cereals (eg, potassium) may lower blood pressure, phytoestrogens may improve lipid levels and insulin sensitivity, and other levels or possess antioxidant properties. Slowing of starch digestion or absorption and promotion of satiety are possible mechanisms by which whole-grain cereals may help control body weight. 109

risk, with about a 20% lower risk associated with an intake of disease and premature death in the United States. Tuxedos one or two times per week and about a 30% lower risk with intake of three or more times per week, compared with intake

less than one time per month. Estimated intake of marine n-3 fatty acids was associated with 37% lower heart failure risk in the highest quintile of intake compared with the lowest. 111 Fish oil favorably affects hemodynamics and reduces blood pressure, inflammation, vascular responses, and myocardial oxygen consumption at given workloads; it increases contractile recovery after ischemia-reperfusion, augments left ventricular response to exercise, prevents left ventricular remodeling, and improves left ventricular indices and diastolic filling.

Short-term trials of fish oil supplementation of 3 to 5 g/day may also reduce risk, whereas dietary doses of about 0.5 g/day may result in more modest effects that during the long term may reduce heart failure risk. The beneficial associations were most pronounced 10 Initially, this diet was promoted for hypertension; however, among persons consuming broiled or baked fish at least three times per week, the equivalent of about 500 mg/day of eicosapentaenoic of heart failure risk, with an observed 37% lower heart failure acid and docosahexaenoic acid; fried fish intake does not seem to exert this benefit on heart failure risk. It has been reported that broiled or baked fish consumption is inversely associated with systemic blood pressure, C-reactive protein levels, and carotid intimal medial thickness, whereas fried fish intake is positively associated with them, indicating that the type of cooking could have an impact on the effects. 111

Historically, human ancestors consumed less than 0.25 g of salt per day; humans may therefore be genetically programmed to this statistically significant associations with cardiovascular amount of salt. 117 The recent change to the high-salt intake of 10 to events. However, women with diets most consistent with 12 g/day presents a major challenge to the physiological systems to excrete these large amounts of salt, resulting in a rise in blood pressure, increase in the risk for cardiovascular and renal disease, bone demineralization, and stomach cancer. 118 The Department of Significantly, the DASH diet reduces low-density lipoprotein Health and Human Services and the Department of Agriculture currently recommend that adults consume no more than 2300 mg/day of sodium (equal to approximately 1 tablespoon of salt), but specific groups (ie, all persons with hypertension, all middleaged and older adults, and all blacks) should consume no more than diet and heart failure has been investigated in human and 1500 mg/day of sodium. 119 Overall, 69.2% of US adults, animal studies. In prospective studies of free-living approximately 145.5 million persons, meet the criteria for the risk groups. 120

There is overwhelming evidence for a causal relationship between salt intake and blood pressure from epidemiology, intervention, treatment, animal, and genetic studies. 106,117,118 Salt intake is also associated with increased risk for overall cardiovascular diseases. A reduction in salt intake may have other beneficial effects on the cardiovascular system, independent of and additive to its effect on blood pressure, including regression of left ventricular hypertrophy, delay in deterioration of renal function, and reduction in proteinuria. 121,122 Salt intake is also associated with incident heart failure in overweight individuals. A high dietary intake of sodium could lead to heart failure because of increased blood pressure (pressure overload) or extracellular fluid (volume overload) and left ventricular hypertrophy.

Specific strategies should be implemented to target an intake of and high-fat diets were compared, animals fed the high- 1500 mg/day of sodium. The food industry should be encouraged fructose diet demonstrated more cardiac remodeling and to reduce sodium used for food preparation. A public health campaign to educate consumers about the dangers of high salt intake and the need to make wise and healthy choices in the sodium content of their foods is of paramount importance. If consumers become more attuned to the sodium content of their foods and the detrimental health effects of a high-salt intake, they could demand low-sodium products, encouraging manufacturers to reduce the sodium content of their products, and they could also prepare their constituents exert beneficial effects on lipid and homocystine food with less salt. Finally, the responsibility to use fresh products and to avoid canned and other high-salt food eventually falls on individuals.

Fish consumption exerts beneficial effects on heart failure Tobacco use is the single most prevalent preventable cause of

is a strong and independent predictor of incident heart failure in both men and women, with 45% and 88% increased risk, respectively, after adjustment for coronary heart disease, implying a more direct effect of smoking on the development of heart failure. ²⁰ In the Coronary Artery Surgery Study, current smokers had a 47% mortality. In the Studies of Left Ventricular Dysfunction trials, exhigher risk for heart failure. 123 Similarly, a cohort study in Sweden smokers had a 30% lower mortality than that of current smokers, a showed that smokers have a 60% higher risk for heart failure, 38 and benefit that accrued within 2 years after smoking cessation. This in the Health ABC Study, current smokers had a twofold higher risk survival rate was similar to that of nonsmokers; the risk for heart for heart failure. 17

Nutrition Examination Survey Epidemiological Follow-up Study ²⁰ and 5.5% in whites and 15% in blacks as reported in the Health ABC Study. 17 The deleterious effect of tobacco seems to be independent **Obstructive Sleep Apnea** of the form of use (smoked or not); increased risk for cardiovascular diseases is reported in nonsmoking use of tobacco. 124 There is no "safe" level of smoking; a single cigarette may stiffen the left ventricle, ¹²⁵ and as few as one to four cigarettes a day double the risk of having a myocardial infarction. 126 Moreover, cigarettes with lower yields of tar and nicotine have not been shown to lower risk of heart disease and should not be considered lower risk alternatives to regular cigarettes.

Mechanisms leading to heart failure in smokers include indirect effects (ie, by causing or aggravating comorbidities that are strongly related to heart failure) and direct effects on the myocardium. Smoking is associated with coronary vasoconstriction, abnormal coronary endothelial function, increased ischemic burden, oxidative hypertension 147 and worsening blood pressure control, 148 coronary stress, increased peripheral vascular resistance, insulin resistance, and type 2 diabetes. 127-130 Cigarette smoke contains superoxide and other reactive oxygen species, 131 which could cause oxidative of hypertension 151,152 and in established heart failure results in damage in endothelial cells ¹³² and cell death. ¹³³ Acute inhalation of nicotine decreases nitrate, nitrite, and serum antioxidant concentrations in the plasma 134 and increases arterial stiffness. 135 failure and was associated with a 2.38 relative risk of heart failure Cigarettes with high- or low-nicotine content or nicotine-free when the apnea-hypopnea index was > 11 events per hour. cigarettes composed of synthetic material increase blood carboxyhemoglobin levels, which in turn decreases the amount of oxygen available to the myocardium. In animal models, nicotine heart failure and (2) neurohormonal abnormalities and mechanical exposure induces interstitial fibrosis in the ventricles. 136 Besides modifications. Sleep is normally a period of cardiac rest. Apart from nicotine, carbon monoxide is also a significant component of tobacco smoke and causes overexpression of growth-related proteins (such as calmodulin, calcineurin, and vascular endothelial increased vagal activity, which lowers heart rate and blood growth factor), impaired cardiac contraction-relaxation cycle, impairment in calcium handling associated with the dysfunction of the SERCA-2a calcium pump, increased cardiomyocyte cGMP, and reduction of T-tubule density, which in turn decreases the synchrony of activation and reduces the rate of calcium release during systole. 137 In healthy humans, smoking is associated with higher left ventricular mass, lower stroke volume, and lower ejection fraction 138 and impaired ventricular diastolic function. 139 All these affected pathways under score the impact of smoking on myocardial status.

Interventions. All individuals should be asked about tobacco use, and smokers should be counseled to quit. Patients should be referred to formal cessation programs, and pharmacological therapy should be offered to increase the success rate. Current recommended strategies include the following.

- · Medications. Several effective medications are available for tobacco addiction, and clinicians should encourage their use by all Hypertension patients attempting to quit smoking. Seven first-line medications Hypertension is an antecedent condition in the majority of reliably increase long-term smoking abstinence rates, including individuals developing heart failure. 38 Systolic blood bupropion SR, nicotine gum or inhaler or lozenge or nasal spray or patch, and varenicline. Notably, none of these medications is contraindicated if cardiovascular disease exists; however, nicotine replacement therapy should be used with caution among particularly cardiovascular patient groups.
- Counseling and psychosocial support. Individual, group, and telephone practical counseling (problem solving, skills training) and social support are effective, and their effectiveness increases with treatment intensity.
- Combination. The combination of counseling and medication,

however, is more effective than either alone. There fore, clinicians should encourage all individuals making a quit attempt to use both counseling and medication.

Quitting tobacco is associated with reduced morbidity and failure hospitalizations and myocardial infarctions also reduced Smoking is carrying a considerable population attributable risk after quitting. 140 Women's risk of heart disease is reduced by one for heart failure: 17% as reported in the First National Health and third within 2 years of quitting and by about two thirds within 5



Obstructive sleep apnea is characterized by abnormal collapse of the pharyngeal airway during sleep, causing repeated arousals. Obesity is a major risk factor for this condition, partly because layering of fat adjacent to the pharynx narrows its lumen. 142 The Wisconsin Sleep Cohort Study, ¹⁴³ a large population-based study, reported that obstructive sleep apnea affects approximately 15% of men and 5% of women between the ages of 30 and 60 years when sleep apnea is defined as an apnea-hypopnea index of > 10 events per hour. Other studies reported similar findings 144,145 or an even higher prevalence, especially in women. 146 Several well-conducted studies provided compelling evidence that obstructive sleep apnea is related to artery disease, 149 and diabetes. 150 In addition, obstructive sleep apnea seems to induce left ven tricular dysfunction independently worse outcomes. 153 In the Sleep Heart Health Study, obstructive sleep apnea was found to be an independent risk factor for heart

Obstructive sleep apnea may lead to heart failure through (1) development and worsening of comorbidities that predispose to brief bursts of sympathetic activity in rapid eye movement (REM) sleep, sleep is characterized by decreased sympathetic and pressure. In patients with obstructive sleep apnea, however, negative intrathoracic pressure generated by the inspiratory effort during obstructed breath increases left ventricular afterload, changes venous return affecting preload and stroke volume, and increases cardiac muscle work index. In addition, sympathetic activation secondary to hypoxia increases blood pressure and heart rate, thus increasing cardiac afterload, and reduces myocardial perfusion, leading to myocardial ischemia. On the other hand, the increased venous return accompanied by acute hypoxic pulmonary vasoconstriction increases right ventricular volume and pressure and may also compromise left ven tricular filling. 154 Sleep apnea treatment with devices that provide continuous positive airway pressure has been shown to improve left ventricular structure and function in patients with established left ventricular dysfunction ¹⁵⁵ or to reverse functional ventricular abnormalities. 151



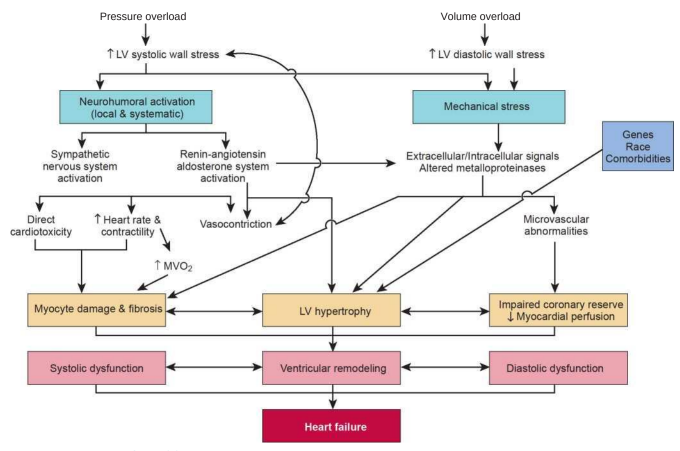


FIGURE 10-6 Hypertension and heart failure risk. LV, left ventricle; MVO2, myocardial oxygen consumption.

pressure increases almost linearly with age, as does the overall lifestyle, 162 the importance of hypertension in the development of prevalence of hypertension and the proportion of patients with heart failure cannot be overemphasized. isolated systolic hypertension. By the age of 75 years, almost all hypertensive individuals have isolated systolic hypertension. 156 changes and systolic and diastolic ventricular dysfunction is well Diastolic hypertension is more prevalent before the age of 50 years, established; Figure 10-6 summarizes this process. Increases in whereas prevalent systolic hypertension increases with age and cardiac afterload, left ventricular mass, and wall stress accompanied after the age of 50 years represents the most common form of by impairment of diastolic filling proper ties occur in the chronic hypertension. 156 Diastolic blood pressure is a more potent setting. Increased peripheral vascular resistance places a greater cardiovascular risk factor than systolic blood pressure until the age myocardial burden leading to an increase in myocardial muscle of 50 years, and thereafter systolic blood pressure becomes more mass. 163 The development of hypertrophy is associated with important. 157 Clinical trials have demonstrated that controlling progressive degenerative changes in the myocytes and an abnormal systolic hypertension reduces heart failure rates. 158 The population accumulation of collagen in the interstitial spaces. 164 Initially, this attributable risk of hypertension for heart failure in the general leads to diastolic dysfunction with increased myocardial stiffness. population is reported to be 39% in men and 59% in women by the 165 Also, the disproportionately increased left ventricular mass leads Framingham investigators, 159 whereas the population attributable to inadequate microvasculature to perfuse the hypertrophied risk of uncontrolled blood pressure in the elderly was reported to be myocardium, resulting in subendocardial hypo perfusion and 21.3% in whites to 30.1% in blacks in the Health ABC Study. ¹⁷ This risk increases in a continuous fashion with increase in blood

pressure.

The lifetime risk for heart failure doubles in subjects with blood pressure > 160/100 mm Hg versus those with < 140/90 mm Hg, and this gradient of risk is seen in both sexes in every decade from 40 to 70 years. 160 Considering that the prevalence of hypertension is estimated to range from 25% to 60% 161 and that this proportion will likely increase with aging of the population and a sedentary

The progression from hypertension to structural ventricular ischemia. 166 Hypertension also contributes to ischemia by increasing myocardial oxygen demand due to increased workload ¹⁶⁷ and is associated with endothelial dysfunction, oxidative stress, and development of atherosclerosis. These changes increase the risk for coronary thrombosis and myocardial infarction characterized by loss of contractile function, neurohormonal activation, and ventricular remodeling, leading to the development of systolic dysfunction. 168

Abnormalities in the neurohormonal activation and water and electrolyte balance also play a central role in the cascade

that leads from hypertension to heart failure. 169 The reninangiotensin-aldosterone system activity increases hypertrophy 169 and heart failure. Angiotensin II is an important initiator of extracellular matrix remodeling, ^{169,170} which contributes to the pathogenesis of atherosclerosis and cardiac hypertrophy. 169 The heightened sympathetic nervous system predisposes to vasoconstriction, sodium retention, and ven tricular hypertrophy. The ventricular hypertrophy occurs as increased norepinephrine release results in myocyte hypertrophy, increased apoptosis of cardiomyocytes, and deficits in cardiomyocyte contractility. 169 These changes are facilitated by beta-adrenergic receptor hyperactivation.

Intervention. The placebo-controlled trials and the meta analysis arising from them demonstrate the benefit of antihypertensive therapy in reducing the incidence of cardiovascular diseases. 171 Fewer studies have specifically focused on prevention of left ventricular hypertrophy and development of heart failure. Systolic Hypertension in the Elderly Program (SHEP) demonstrated that antihypertensive treatment compared with placebo exerted a strong protective effect, ¹⁵⁸ whereas a meta-analysis of 12 hypertension trials that included the development of heart failure and 4 that included the incidence of left ventricular hypertrophy as endpoints demonstrated significant treatment benefits. 172 The incidence of left ventricular hypertrophy was decreased by 35% (95% CI, 21% to 48%) and the incidence of heart failure was reduced by 52% (95% CI, 41% to 62%) compared with placebo subjects (Fig. 10-7).

Antihypertensive Medications

· Diuretics. Thiazide diuretics have been effective in preventing cardiovascular complications of hypertension. Secondary outcomes of the Antihypertensive and Lipid-Lowering Treatment on the effects of calcium antagonists on left ventricular mass or to Prevent Heart Attack Trial reported a higher rate of incident incident heart failure. A recent meta-analysis suggests that heart failure with amlodipine (relative risk of 1.35) and a nonsignificant increase with lisinopril (relative risk of 1.09) compared with chlorthali done. 173 On the contrary, the Second Australian National Blood Pressure trial 174 reported better outcomes with a regimen that was initiated with an angiotensinconverting enzyme inhibitor compared with a diuretic. Diuretics are at least as good as other classes of drugs and also enhance the antihypertensive efficacy of multidrug regimens; the Joint National Commission 7 recommends that in the absence of any other compelling indications, thiazides

diuretics should be used as initial therapy for **153** hypertension. Renin-angiotensin system modulators. Meta-analysis of double-blind trials that measured the effects of antihypertensive drugs on left ventricular mass 175 showed that the greatest reduction was

achieved with angiotensin receptor blockers (Fig. 10-8). Also, these agents along with calcium channel blockers and angiotensinconverting enzyme inhibitors were more effective than beta blockers in reducing left ventricular mass. Both irbesartan- and losartan-based regimens have been shown to reduce left ventricular I mass more than atenolol-based treatments. 176,177 Candesar tan had efficacy similar to that of enalapril. 178 These data suggest that antihypertensive agents targeting the renin angiotensin system are particularly effective in producing 'Q regression of left ventricular hypertrophy and that angio tensin receptor blockers are at least as effective as angiotensin-converting enzyme inhibitors. A recent meta analysis of renin-angiotensin system inhibition showed that these agents reduce the risk of heart failure by 19% compared with calcium channel blockers. 179

Beta blockers. Although beta blockers are effective in lowering blood pressure, these medications are less effective in preventing hypertension complications, including coronary artery disease and cardiovascular and all-cause mortality, ^{180,181} or in reducing left ventricular mass. ^{176,177} However, a recent meta-analysis suggested that beta blockers are effective for primary prevention of heart failure in hypertension and had a similar benefit in the elderly and younger individuals compared with other agents. 182 On the other hand, the 19% increased risk for stroke in the elderly associated with beta blocker use in the same analysis tempers the enthusiasm to use them as first-line agents for heart failure prevention in hypertension. 182

Calcium channel blockers. There are limited experimental data treatment of hypertension with calcium channel blockers is less effective for reducing heart failure for the same reduction of blood pressure 179; the effect of this class of medications on reducing left ventricular mass was similar to that of reninangiotensin system inhibition. 175 Target Goals of Therapy. In patients with hypertension, systolic and diastolic blood pressure targets are

154 < 140/90 mm Hg except for patients with diabetes or renal disease, for whom the goal is < 130/80 mm Hg. Because most patients with hypertension, especially those older than 50 years, will reach the diastolic blood pressure goal once systolic blood pressure is at goal, the primary focus should therefore be on systolic blood pressure. Current control rates at the population level remain far below the Healthy People 2010 goal of 50%,

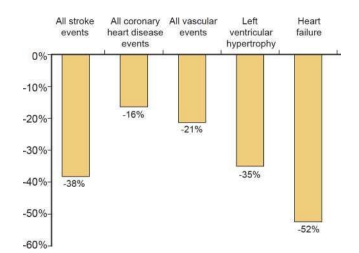
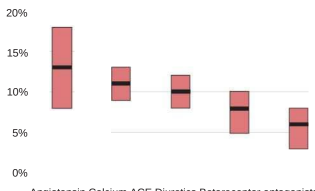


FIGURE 10-7 Hypertension treatment and its impact on cardiovascular events. Results of meta-analyses of randomized trials of antihypertensive medications therapy, indicating the impact of blood pressure reduction on reduction of cardiovascular, vascular, heart failure, and left ventricular hypertrophy events. 71.72



Angiotensin Calcium ACE Diuretics Betareceptor antagonists inhibitors blockers

FIGURE 10-8 Alteration in left ventricular mass index with various antihypertensive medications. Angiotensin II receptor antagonists, calcium antagonists, and angiotensinconverting enzyme (ACE) inhibitors reduced left ventricular mass index more than beta blockers did. (Reproduced from Klingbeil AU, Schneider M, Martus P, et al: A meta-analysis of the effects of treatment on left ventricular mass in essential hypertension. Am J Med 115:41, 2003.)

with 30% of patients still being unaware of their hypertension status. Recent trials have demonstrated that effective blood pressure control can be achieved in most patients, but the majority will require two or more medications in combination. 183 Data from a recent randomized trial evaluating the effect of usual versus tight control of systolic blood pressure (< 130 mm Hg) in nondiabetic hypertensive 10 individuals with left ventricular hypertrophy demonstrated additional benefit with tighter control. In particular, left ven tricular hypertrophy was less frequent, but also the composite outcome of all-cause mortality, cardiovascular events, and heart failure was lower in the tight control group. 184

Diabetes Mellitus

Diabetes mellitus is an independent risk factor for the development of heart failure in all age groups. 20,37 The relative risk for new-onset heart failure among patients with diabetes mellitus ranges from 1.3 to 2.7, increasing to 4 in patients younger than 65 years and 11 in those younger than 45 years specifically.

Several mechanisms have been proposed to explain the increased risk for heart failure among patients with diabetes mellitus. Comorbidities associated with heart failure, including obesity, hypertension, and coronary artery disease, are highly prevalent among individuals with diabetes mellitus. Insulin resistance itself may produce abnormalities in cardiac structure and function. 185 Patients with insulin resistance exhibit endothelial dysfunction proinflammatory state, which contribute to ventricular dysfunction, even before the development of overt diabetes mellitus. Left ventricular hypertrophy and left ventricular dysfunction are also strongly correlated with insulin resistance, and hyperinsulinemia has been associated with sympathetic nervous system activation. 185

Several mechanisms have been proposed for development of "diabetic cardiomyopathy" 186,187:

- Microangiopathy and endothelial dysfunction.
- Autonomic neuropathy, which causes impaired coronary vasodilatory response to sympathetic stimulation and modulates the contractility of cardiac myocytes.
- Metabolic derangements caused by insulin resistance, such as reduced glucose and lactate metabolism and enhanced fatty acid metabolism, result in lipid accumulation in the myocardium, promoting lipotoxicity. This increased fatty acid metabolism causes increased myocardial oxygen consumption, promoting ischemia and arrhythmias, inflammatory pathways, enhance smooth muscle cell proliferation, and contribute to endothelial dysfunction. ¹⁸⁸
- Abnormalities in ion homeostasis through alteration of ion alteration of the function or cardiac expression of sodium calcium exchangers, sarcoplasmic reticulum/sarcolemmal patients with diabetes. 193 calcium-ATPases (SERCA-2a), calcium-binding proteins, and the mitochondrial calcium uniporter.
- Upregulation of the renin-angiotensin system.
- Increased oxidative stress.
- Increased glycation of interstitial proteins such as collagen, creating advanced glycosylation end products, which result in myocardial stiffness and impaired contractility.
- Activation of protein kinase C leads to alterations in cell growth and function, affecting muscle contractility and gene expression.

Interventions

Medications

Insulin. Whether insulin use specifically reduces the risk of heart failure is not known. Randomized controlled trials indicate that insulin use in ACC/AHA stage A heart failure does not appear to significantly increase the risk of new-onset heart failure, 189 and insulin use in stage B heart failure does not appear to negatively affect mortality. 188

- Sulfonylureas. The available data suggest that sulfonylurea therapy does not significantly increase the risk of heart failure compared with other oral antidiabetic agents, although the ADOPT (A Diabetes Outcome Progression Trial) was not powered to detect significant differences. 190
- Metformin. The risk of new-onset heart failure among patients treated with metformin compared with patients treated with other oral antidiabetic medications was reported in ADOPT, and the findings were similar to those for sulfonylureas. 190
- Thiazolidinediones. The hypothesis that therapies that will correct abnormal myocardial substrate metabolism in diabetes will translate to a lower incidence of heart failure appears promising; however, the available data on this class of medications argue hypothesis. Although treatment thiazolidinediones increased myocardial glucose uptake in patients with underlying coronary disease, and myocardial glucose uptake seems to be positively correlated with left ventricular function, 188 the Rosiglitazone Evaluated for Cardiac Outcomes and Regulation of Glycaemia in Diabetes study has indicated that thiazolidinedione use is associated with a small but clinically relevant increased risk of heart failure in patients with ACC/AHA stage A or stage B status. 34

Other agents. There are limited data regarding the risks or benefits of other antidiabetic therapies for incident heart failure, except for the a -glucosidase inhibitor acarbose, the impact of which on cardiovascular disease outcomes was evaluated in the STOP-Noninsulin-Dependent Diabetes Mellitus (STOP-NIDDM) trial. 191 Although the study was not powered to evaluate the impact of acarbose treatment on the development of heart failure, the available evidence suggests that this agent may decrease the risk of myocardial infarction and subsequent risk of heart failure. 191

Glucose Control. The best available evidence suggests that management of hyperglycemia in patients with type 2 diabetes mellitus reduces neither cardiovascular morbidity and mortality nor the progression of heart failure. In the United Kingdom Prospective Diabetes Study (UKPDS) 33, no significant reduction in the development of macrovascular disease or heart failure was demonstrated with intensive blood glucose control. 189

Blood Pressure Control in Diabetes Mellitus. Because hypertension further increases the risk of cardiovascular disease and heart failure in patients with diabetes, aggressive blood pressure management is essential to prevent long-term complications in this population. In UKPDS 38, tight blood pressure control with either the angiotensin-converting enzyme inhibitor or beta blocker whereas lipid intermediates (eg, ceramide) might promote significantly reduced the risk of cardiovascular events and diabetesapoptosis. Elevated fatty acids also activate coagulation and related mortality, including a 56% reduction in the risk for heart failure. ¹⁹² Notably, target blood pressure level was < 150/85 mm Hg in that study, whereas the Seventh Report of the Joint National Commission on Prevention, Detection, Evaluation, and Treat ment channels such as calcium and potassium channels and of High Blood Pressure (JNC 7) recommended more aggressive blood pressure control (target blood pressure < 130/80 mm Hg) in

> Targeting Mechanisms Causing Diabetic Cardiomyopa thy. The mechanisms leading to diabetic cardiomyopathy are

inhibitors, angiotensin receptor blockers, and beta blockers benefit modulate the reparative changes, including dilation, hypertrophy, including heart failure. The Heart Outcomes Prevention Evaluation and oxidative stress. 206 (HOPE) trial 194 and the Microalbuminuria, Cardiovascular, and Renal Outcomes (MICRO-HOPE), a sub study of HOPE, 195 have shown that treatment with angiotensin converting enzyme inhibitors reduces the relative risk for new-onset heart failure by 23% and 20%, respectively, in a population of ACC/AHA stage A or B patients, whereas the extension of HOPE study (extended follow-up period) revealed that the benefit of angiotensin-converting enzyme inhibitors for heart failure prevention is sustained over time. 196 Likewise, the Reduction of Endpoints in Non-Insulin Dependent Diabetes Mellitus with the Angiotensin II Antagonist Losartan (RENAAL) study 197 and the Losartan Intervention for Endpoint Reductions in Hypertension (LIFE) study 198 have shown a 32% and 41% reduction in the frequency of hospitalization for heart failure, respectively.

A recent meta-analysis evaluating the beneficial effects of the for new-onset heart failure showed that their favorable effect was that stimulates hypertrophy mediated by mechanoreceptors and peripheral edema, which is common among patients receiving These changes result in alterations in circulatory hemodynamics, calcium channel blockers and may complicate the diagnosis of heart failure. Recently, a randomized trial evaluating the effects of the restoration of left ventricular mass towards normal. In recent studies, deformation. breakers of advanced glycation end products-related protein crossassociated with aging, diabetes, and hypertension. 200 These effects support the hypothesis that modulation of specific pathways could

Coronary Heart Disease

Coronary heart disease is a major antecedent condition predisposing to an increased risk for development of heart failure in both men and Chronic Coronary Artery Disease women, 44 with reported population attributable risks ranging from Ischemia caused by abnormalities in coronary arteries can produce 56% in women to 62% in men by the First National Health and increases in the concentration of neurohormones Nutrition Examination Survey Epidemiological Follow-up Study 20 norepinephrine, epinephrine, endothelin, and dopa mine) that and from 23.9% in whites to 29.5% in blacks as reported in the Health results in myocardial apoptosis, fibrosis, and susceptibility to ABC Study. 17 Heart failure incidence in the Framingham Heart ventricular arrhythmias. 212 Thus, ischemia contributes to the Study changed only modestly from the 1950s through the 1990s in progression of left ventricular systolic dysfunction even in the men but declined by about one third among women during this absence of a manifest infarct event. Chronic ischemia can result in period. This trend was also supported by other cohort studies. 201,202 hibernation or stunning with further progressive decline in These trends in men and women may be partially attributed to sex-ventricular function. These adaptive-protective mechanisms may based differences in heart failure etiology because hypertension result in myocardium rendered hypocontractile and contribute to predominates as an etiologic risk factor for heart failure in women overall left ventricular systolic dysfunction. Hibernation represents more so than in men, and coronary heart disease is responsible for a a pre- carious balance between perfusion and tissue viability that higher proportion of cases in men. 20,159 Increased awareness and cannot be maintained indefinitely, and myocardial necrosis will treatment of hypertension have led to improved trends in control 203 eventually occur if blood flow is not increased. 207,212 Must and probably decreased heart failure incidence more so in women, especially because blood pressure control appears to be better in women. ²⁰⁴ On the contrary, advances in the treatment of myocardial infarction ²⁰⁵ have led to increasing numbers of patients surviving with residual myocardial damage, which may be partially responsible for the increase in heart failure incidence among men. 20

Acute Myocardial Infarction

Acute myocardial infarction leads to a cascade of adaptive mechanisms that promote left ventricular remodeling. Acute loss of myocardial cells results in an increase in loading conditions on the remaining myocardium and induces a unique pattern of remodeling involving the infarcted border zone and the noninfarcted

complex. Medications like angiotensin-converting enzyme myocardium. Intracellular signaling processes initiate and 155 patients with diabetes and prevent complications of diabetes formation of a collagen scar, neurohormonal and cytokine activation,

> Ventricular remodeling may continue for weeks or months until the distending forces are counterbalanced by the tensile strength of the collagen scar; this balance then determines the size, location, and transmurality of the infarct and the extent of myocardial stunning, ventricular loading conditions, and local trophic factors. 207 Although contemporary treatment attenuates remodeling, ^{208,209} there is a large heterogeneity in the remodeling response after an infarction. Whereas remodeling in the short term is adaptive, over time it becomes deleterious, leading to adverse structural and hemodynamic changes leading to heart failure. 209

Initially, myocardial infarction results in the migration of circulating inflammatory cells into the infarct zone, triggering 10 intracellular signaling and neurohormonal activation, leading to a local inflammatory response that in turn leads to cell death and degradation of the intermyocyte collagen struts, expanding infarct size. 210 Infarct expansion results in wall thinning and ventricular agents that inhibit the renin-angiotensin system on reducing the risk dilation and causes the elevation of diastolic and systolic wall stress beyond blood pressure control, and the risk was 19% lower than for transduces intracellular signaling, which initiates the increased calcium channel blockers. 179 Although these data are promising, synthesis of contractile assembly units. 210 Infarct expansion causes there are concerns about the heterogeneity of the criteria used for deformation of the border zone and remote myocardium, which assessment of heart failure in the various trials and the side effect of alters the Frank-Starling response and augments shortening. 210 triggering sympathetic adrenergic and renin-angiotensinaldosterone systems, 206 leading to myocyte hypertrophy and Cu(II)-selective chelator trientine on left ventricular hypertrophy in alterations in the ventricular architecture to distribute the increased diabetes mellitus showed a 10% decrease in left ventricular mass wall stress more evenly as the extracellular matrix forms a collagen index after 12 months. 199 This decrease represented a ~50% scar to stabilize the distending forces and to prevent further

Reperfusion therapy helps prevent infarct progression. However, links improved the adverse cardiovascular and renal changes it is associated with generation of reactive oxygen species, osmotic gradient and cell swelling, activation of the sodium-hydrogen exchanger, calcium overload-induced myocardial contracture, local lead to attenuation of diabetes complications including heart failure. inflammatory and oxidant response to reperfusion, and opening of the mitochondrial permeability transition pore that extends infarct size beyond that observed during equivalent periods of ischemia alone. 211 Thus, reperfusion injury is a possible target for interventions to reduce myocardial damage.

156 patients with heart failure of ischemic origin have a substantial volume of myocardium that fails to contract because it is stunned or hibernating rather than because it is scarred. In addition, endothelial dysfunction, an inherent component of the pathophysiological process of atherosclerosis, could directly affect ventricular function. 213 Ischemic mitral regurgitation, caused by changes in ventricular structure and function, increases left ventricular preload, leading finally to alteration of left ventricle geometry and deterioration of its function. 214

Moreover, myocardial ischemia induces dysfunction related to impaired calcium ion sequestration into the sarcoplasmic reticulum during the energy-dependent process of therapy. 212

myocyte reserve; however, in patients with established ventricular remodeling can be reversed. ²²⁸ coronary heart disease, aggressive management can reduce the the risk of heart failure.

for myocardial salvage. In addition, the "open artery Survival Study (EPHESUS). 229 hypothesis" proposes that late reperfusion, beyond the their younger counterparts. 219

Angiotensin-Converting Enzyme Inhibitors. Angiotensin -²²⁰ Similar findings were demonstrated in the Survival of Myocardial Infarction Long-term Evaluation trial with zofenovascular disease, 194 expanding the indication for angiotensindocumented coronary heart disease, presumed coronary heart failure and to decrease mortality in this population. disease based on the presence of other atherosclerotic vascular disease, or diabetes. The European trial on Reduction of insights into cardiac myocyte function, repair cardiac events with Perindopril in stable coronary artery disease (EUROPE) showed similar data with per indopril. 222 In the Survival and Ventricular Enlargement (SAVE) trial that enrolled patients with asymptomatic left ventricular dysfunction, captopril led to a 22% reduction in the risk of heart failure hospitalization. 223

Angiotensin Receptor Blockers. Available data suggest that angiotensin receptor blockers are at least as effective as angiotensinconverting enzyme inhibitors in reducing mortality in patients with myocardial infarction complicated by left ventricular dysfunction or heart failure. 207,212 However, data on populations with atherosclerotic diseases but without heart failure are not uniform. ^{224,225} Because the overall evidence for the effectiveness of angiotensin receptor blockers on prevention and attenuation of post-myocardial infarction left ventricular remodeling is weaker compared with angiotensin-converting enzyme inhibitors, they may not be used as first-line therapy but limited to those individuals who do not tolerate angiotensin-converting enzyme inhibitors.

Beta blockers. Beta blockers have been proven to be effective after relaxation 215 and through alteration of the myocardial 10 passive acute myocardial infarction for more than 20 years. Notably, longcompliance resulting from scarring, fibrosis, and compensatory term beta blocker use is recommended for secondary prevention in hypertrophy of noninfarcted myocardium. ²¹⁶ Diastolic patients at highest risk (eg, those with low ejection fraction or heart dysfunction during myocardial ischemia precedes systolic failure). ²²⁶ The echo cardiographic substudy of Carvedilol Postdysfunction and takes longer to recover from. 212 Left ventricular Infarct Survival Control in Left Ventricular Dysfunction diastolic dysfunction is present in the very early phases of (CAPRICORN) demonstrated a beneficial effect of carvedilol on left myocardial infarction, and it is associated with the development of ventricular remodeling in patients with left ventricular dysfunction in-hospital heart failure and cardiac death; several studies have after myocardial infarction over that of angiotensin-converting documented recovery of diastolic function after reperfusion enzyme inhibitors. 227 The Reversal of Ventricular Remodeling with Toprol-XL (REVERT) trial gives further evidence that beta blocker Interventions. Prevention of coronary heart disease and use for the treatment of asymptomatic left ven tricular dysfunction ischemic events is a key point to maintenance of functional to prevent development of heart failure is effective and that left

Aldosterone Antagonists. High aldosterone levels seen in development of heart failure. A number of cardioprotective patients with myocardial infarction induce ventricular remodeling. medications and procedures can prevent the development of ²⁰⁷ Spironolactone combined with angiotensin converting enzyme symptomatic heart failure in coronary heart disease. The inhibitors improved left ventricle remodeling after acute myocardial combination of medications along with therapeutic lifestyle infarction better than angiotensin-converting enzyme inhibitors changes as recommended in the American Heart Association/ alone did in a randomized study. 207 This study also showed that American College of Cardiology secondary prevention guide spironolactone significantly suppressed transcardiac extraction of lines 217 should be applied aggressively in all patients to reduce aldosterone and plasma levels of procollagen type III amino-terminal peptide, a marker of cardiac fibrosis. Aldosterone antagonists are Revascularization. Mechanical (percutaneous or surgical) recommended in myocardial infarction complicated by left or pharmacological revascularization of the infarct-related ventricular dysfunction on the basis of the beneficial effect on artery reduces the size of the acute infarct and prevents - mortality and cardiovascular hospitalizations seen in the Eplerenone subsequent heart failure 208,210 if it is performed early enough Post-Acute Myocardial Infarction Heart Failure Efficacy and

Antiplatelet Agents. Aspirin in patients with established vascular window for myocardial salvage, also reduces left ventricular disease has been demonstrated to reduce the risk of cardiovascular remodeling. ²¹⁸ Even after successful early revascularization, events and heart failure. ²¹² It is recommended after acute myocardial older patients are at higher risk for heart failure compared with infarction and should be continued indefinitely if no contraindications exist. 230

Statins. Statins are of proven benefit in patients with coronary converting enzyme inhibitors have favorable properties in heart disease 212,231; however, their usefulness in the setting of left reducing left ventricular stress and progression of left ven - ventricular dysfunction remains under investigation. Preprocedural tricle enlargement, and several studies have shown them to treatment with a statin before percutaneous coronary intervention is reduce the incidence of heart failure and mortality after an associated with lower levels of periprocedural creatine kinase acute myocardial infarction. In the third Gruppo Italiano per elevation. 207 Ishii and coworkers 232 reported that chronic statin lo Studio della Sopravvivenza nell'Infarto Miocardico (GISSI) therapy before the onset of the acute event is associated with trial, early lisinopril therapy in acute infarction reduced improved perfusion and reduced myocardial necrosis after the mortality and left ventricular dysfunction despite therapy with intervention. Prospective studies in humans reported an effect of aspirin, thrombolytics, and beta-adrenergic blocking agents. statins on the risk for development of heart failure in high-risk patients, most of whom were free of heart failure at enrollment. Kjekshus and associates ²³³ showed an 11% lower risk of new-onset pril. ²²¹ The HOPE study demonstrated a 23% reduction in risk heart failure in patients with stable coronary heart disease treated for heart failure by ramipril in individuals with established - with statins. Similar trends were also demonstrated in other studies, ²³⁴ supporting a role for statins in the prevention of heart failure. converting enzyme inhibitor therapy to all patients with Notably, statins appear to prevent further progression of heart

Novel Therapeutic Approaches. Several novel mechanistic

mechanisms, and remodeling may lead to the development of newer 8. therapies for heart failure in the future. These include therapies targeted at nitric oxide-cGMP signaling and nitric oxide modulation, oxidative stress and antioxidant therapy, anti-inflammatory therapies, modulation of innate immunity toll-like receptors, 10. Pearson TA, Blair SN, Daniels SR, et al: AHA Guidelines for Primary Prevention of Cardiovascular interleukin-1 receptor antagonists, and selective matrix metalloproteinase inhibition. 231

Stem Cell Therapy. There is also a growing interest in treatment 11. Lloyd-Jones D, Adams R, Carnethon M, et al: Heart disease and stroke statistics—2009 update: a strategies targeting angiogenesis or stem cell transfer. Coronary angiogenesis is enhanced during the acute phase of adaptive cardiac growth but is reduced as maladaptive remodeling progresses. Inhibition of angiogenesis leads to a decreased capillary density, 13. Vasan RS, Benjamin EJ, Levy D: Prevalence, clinical features and prognosis of diastolic heart contractile dysfunction, and impaired cardiac growth. 231 Thus, both cardiac size and function are angiogenesis dependent, and disruption of coordinated tissue growth and angiogenesis in the heart may contribute to the progression from adaptive cardiac hypertrophy to heart failure. Recent observations indicate that stem and progenitor cells can release proangiogenic factors, which in turn stimulate angiogenesis after infarction. Increased angiogenesis after 17. Kalogeropoulos A, Georgiopoulou V, Kritchevsky SB, et al: Epidemiology of incident heart failure stem and progenitor cell transfer has been postulated to improve infarct healing and energy metabolism in the infarct border zone. 231 Early clinical trials suggest that intracoronary delivery of bone marrow cells may improve left ventricular ejection fraction after 19. Butler J. Risk factors for heart failure. In Hosenpud JD, Greenberg BH, editors: Congestive heart infarction. 231 More work is needed, however, to identify the most suitable cell types and application methods and to define the impact 20. He J, Ogden LG, Bazzano LA, et al: Risk factors for congestive heart failure in US men and of cell therapy on clinical endpoints and other indices of ventricular remodeling . Furthermore, other delivery strategies for 21. Fonarow GC: The Acute Decompensated Heart Failure National Registry (ADHERE): opportunities proangiogenic factors after infarction need to be explored.

CONCLUSION

Considering the worsening epidemiological trends and increase in poor quality of life and outcomes for these patients, the importance of heart failure prevention cannot be overemphasized. This will require efforts at all levels of the prevention spectrum ranging from 26. Djousse L, Gaziano JM: Alcohol consumption and risk of heart failure in the Physicians' Health advocacy efforts (eg, to reduce salt in the food chain, population screening of risk factors and their optimal control) to research in understanding novel pathophysiological mechanisms and newer risk factors.

healthy lifestyle habits are essential to promote overall 30. Dhingra R, Sesso HD, Kenchaiah S, et al: Differential effects of lipids on the risk of heart failure cardiovascular health and to reduce heart failure risk specifically. Toward this direction, the American Heart Association has taken a 31. Butler J, Kalogeropoulos A, Georgiopoulou V, et al. Serum resistin concentrations and risk of new bold step in defining their 2020 goals, not only to achieve further reductions in mortality due to cardiovascular disease and stroke but 32. to improve the health of the population based on a comprehensive, specifically designed metric that includes multiple healthy lifestyle 33. Tang WH, Katz R, Brennan ML, et al: Usefulness of myeloperoxidase levels in healthy elderly parameters. Last, whether treatment goals of heart failure risk factors should be individualized on the basis of any given individual's cumulative risk profile needs further study.

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CHAPTER 1 1

Antihypertensive Drugs and Their Cardioprotective and Renoprotective Roles in the Prevention and Management of Cardiovascular Disease

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KEY POINTS

- The complete axis of the RAAS along with intracellular RAAS components, alternative pathways, the kallikrein-kinin system, plasma renin and prorenin, and prorenin receptors play a significant role in cardiovascular and renovascular pathophysiology.
- The relationship between cardiac and renal disease is significant, with a strong but incomplete link between renoprotection and cardiovascular risk reduction.
- Proteinuria may be a potential cardiovascular risk factor and a target or indicator of therapeutic response.
- RAAS inhibitors are the primary classes of antihypertensive medications with both cardioprotective and renoprotective effects and are the cornerstone for prevention, particularly in high-risk patients.
- Combinations of different classes of RAAS inhibitors or RAAS inhibitors with calcium channel blockers provide novel ways for more complete RAAS suppression and improved endorgan protection.

Understanding of the role of the reninangiotensin-aldosterone system (RAAS) in the genesis of cardiac and vascular disease and the close relationship between renal dysfunction and cardiovascular outcomes is increasingly important. Many classes of antihypertensive medications and their combinations have been shown to have beneficial effects, including cardioprotection and renoprotection, and are the cornerstone for primary and secondary prevention. This chapter details each class of antihypertensive medications and elaborates their roles in prevention (Table 11-1).

DIURETICS

Diuretics can be distinguished as loop or thiazide type. Loop-type diuretics have a greater ability to affect volume-overloaded states, and thiazide-type diuretics have a greater ability to reduce blood pressure. Their mechanisms of action differ. Loop diuretics inhibit the sodium-potassium-chloride (Na-K-Cl) transporter on the ascending limb of the loop of Henle. Thia zide diuretics primarily inhibit sodium reabsorption at the renal distal convoluted tubule, thereby reducing plasma volume and lowering blood pressure. Thiazide use also reduces peripheral vascular resistance and acutely stimulates the RAAS. 1-2

Because of the thiazide diuretic role in antihypertensive treatment and its associated outcomes data, thiazides are the main class of diuretics used for primary prevention . The most commonly used thiazide in clinical practice, hydrochlorothiazide, has a short half-life of 8 to 15 hours with long- term dosing. ³ Another thiazide, chlorthali done, has a longer half-life, 45 to 60 hours, and has been used in most clinical trials, establishing the utility of thiazides. ³ Thia zides are particularly beneficial in controlling hypertension in certain populations of salt-sensitive patients, including African Americans, women, and the

elderly. 4

Role of Diuretics

Early literature on diuretics did not show as great a reduction in cardiovascular events as had been predicted by epidemiological data. Many believed that such a discrepancy was explained by the shorter duration of clinical trials compared with epidemiologic studies. Others postulated that the increased risk of cardiovascular effects was attributed to the many potential adverse, metabolic effects of diuretics. which include electrolyte hyperlipidemia, abnormalities, and impaired hyperglycemia, insulin sensitivity. 5-8

Diuretics can alter the lipid profile by increasing both serum low-density lipoprotein (LDL) and very-low-density lipoprotein (VLDL) cholesterol, without having a significant effect on high-density lipoprotein (HDL) cholesterol or the apoproteins. ^{8,9} The mechanism of diuretic-induced dyslipidemia is unclear, and whether this occurs for all patients taking diuretics requires further investigation. Diuretics may also impair insulin sensitivity in both diabetic and non-diabetic patients, which alone or together with resulting compensatory hyperinsulinemia can promote dyslipidemia and atherogenesis. ^{6,8} There may be an increased risk of new-onset diabetes or metabolic syndrome with the use of thiazides, especially at higher doses. ¹⁰

More recent clinical trials question the role that diuretics have in reducing insulin sensitivity. Systolic Hypertension in the Elderly Program (SHEP) and Hypertension in the Very Elderly Trial (HYVET), in which diuretics were used as initial therapy or in combination with other agents, showed a significant reduction in cardiovascular events with no significant change in lipid profiles and a minimal increase in blood glucose levels. ¹¹⁻¹³ A mean follow-up of 14.3

Antihypertensive Drug Class	Examples	Mechanisms of Action	Physiological Effects
Diuretic (thiazide type)	Chlorthalidone Hydrochlorothiazide	Inhibits sodium reabsorption via Na -/Cl - channels in the distal convoluted tubule of the kidney	Plasma volume i Peripheral vascular resistance T RAAS
Nitrates	Nitroglycerin Isosorbide mononitrate Isosorbide dinitrate	Produces active nitric oxide metabolites	T Vasodilation of coronary arteries, collateral vessels, veins Systemic arterial pressure i Preload, myocardial wall stress, and MV o 2 T Endothelial function
Calcium channel blocker	Dihydropyridines: nifedipine, amlodipine, felodipine Nondihydropyridines: diltiazem, verapamil	Reduces excess Ca entry through	T Arteriolar vasodilation including coronary arteries i Inotropy (nondihydropyridine)

			2
			T Endothelial function
Calcium channel blocker	Dihydropyridines: nifedipine, amlodipine, felodipine Nondihydropyridines: diltiazem, verapamil	Reduces excess Ca entry through voltage- and receptor-operated calcium channels in vascular smooth muscle and cardiac myocytes Improved nitric oxide production and release Increases t-PA activity Reduction in A II-mediated vasoconstriction and aldosterone production	T Arteriolar vasodilation including coronary arteries i Inotropy (nondihydropyridine) T Endothelial function T RAAS
Beta blocker	Nonselective beta 1 and beta 2 antagonists: propranolol Selective beta 1 antagonists: atenolol, bisoprolol, metoprolol, nebivolol Alpha 1, beta 1, beta 2 antagonists: carvedilol, labetalol	Decrease in sympathetic activity through blockade of beta 1, beta 2, and alpha 1 receptors Lower plasma renin levels Reduction in endothelin and oxidative stress	In catecholamine levels i Myocardial workload and oxygen demand Fatty acid metabolism i Inotropy, transiently Altered sympathomimetic activity
Angiotensin-converting enzyme (ACE) inhibitor	Benazepril, captopril, enalapril, fosinopril, lisinopril, perindopril, quinapril, ramipril, trandolapril	Inhibits ACE, reducing levels of A II and aldosterone Inhibits ACE-dependent metabolism of bradykinin and kallidin	Alters cardiac remodeling and vascular protective effects Modulation of adrenergic tone T Bradykinin
Angiotensin receptor blocker	Candesartan, irbesartan, losartan, olmesartan, telmisartan	Block the AT 1 receptor Compensatory rise in A II may activate AT 2 receptor	i Vascular and cardiac hypertrophy Aldosterone secretion Vasoconstriction
Direct renin inhibitor	Aliskiren	Blocks renin, the rate-limiting step of A II	i Atherosclerosis progression * and aldosterone secretion i Media degeneration *

production

Reduction in the formation of AI and AII Reduction in plasma renin activity

Blocks aldosterone receptor

Mineralocorticoid receptor blocker

years of patients from the SHEP trial showed that diabetes that developed during the trial among subjects receiving placebo was associated with worsened cardiovascular outcomes. 14 However, diabetes that developed among subjects during diuretic therapy did not significantly affect cardiovascular mortality.

Eplerenone, spironolactone

The Antihypertensive and Lipid-Lowering Treatment to Prevent Heart Attack Trial (ALLHAT) randomized more than 33,000 high-risk hypertensive patients to a regimen of chlorthalidone, amlodipine, or lisinopril as first-line therapy. 15 More than half of the patients enrolled had evidence of atherosclerotic heart disease, with one quarter having a history of myocardial infarction or stroke. No difference in cardio vascular outcomes or mortality was noted among the different agents. Although cholesterol levels and the prevalence of newonset diabetes (11.6% versus 9.8% and 8.1%) were higher in the diuretic group up to a mean 4.9-year follow-up, these metabolic differences did not translate into increased cardiovascular events or mortality. Whether the follow-up period for ALLHAT was adequate to reveal the impact of

dyslipidemia and new-onset diabetes on outcomes remains unclear.

T Nitric oxide availability *

Plasma volume

Cardiac hypertrophy

The SHEP, HYVET, and ALLHAT trials were performed in older populations with an average age of 67 to 84 years. Data from these trials may not be applicable to younger patients with hypertension, particularly in association with comorbid conditions such as obesity, dyslipidemia, and abnormal glucose tolerance. In contrast to ALLHAT, the Second Australian National Blood Pressure (ANBP2) trial showed a significant improvement in cardiovascular outcomes in elderly hypertensive patients who were started with an angiotensinconverting enzyme (ACE) inhibitor compared with a thiazide diuretic. ¹⁶ There were several differences between these trials, including differences in the antihypertensive agents used (enalapril instead of lisinopril and hydrochlorothiazide instead of chlorthalidone) and the population of patients studied (lower baseline risk factors). Nevertheless, thiazide diuretics appear highly effective in lowering blood pressure and reducing cardiovascular events (Table 11-2).

^{*}In animal models

Antihypertensive Drug		
Class Diuretics (thiazide type)	Renal Trials	Renal Effects Unclear effect on proteinuria Potential
	ALLHAT (n = 33,357): chlorthalidone, amlodipine, or lisinopril versus doxazosin; diuretic with significantly increased GFR slope during 4 years compared with CCB (- 0.018 versus - 0.012	worsening of renal function
	dL/mg per year) INSIGHT (n = 6321): nifedipine GITS versus hydrochlorothiazide + amiloride; significant decrease in GFR 2.3 mL/min	
	SHEP (n = 4736): chlorthalidone ± atenolol; significantly increased creatinine 2.8 ^ mol/L	
Nitrates	None	None
Calcium channel blocker (CCB)	AASK (n = 1094): metoprolol, ramipril, or amlodipine; no significant difference in GFR slope; ACE inhibitor showed 38% improvement in composite GFR, ESRD, or death versus CCB ALLHAT (n = 33,357): chlorthalidone, amlodipine, or lisinopril versus doxazosin; CCB with significantly decreased GFR slope during 4 years compared with diuretic (-0.012 versus - 0.018 dL/mg per year) IDNT (n = 1715): irbesartan or amlodipine versus placebo; no significant difference in renal outcomes between amlodipine and placebo	Unclear effect on proteinuria Potential preservation of renal function
Beta blocker		Reduction in proteinuria
	AASK (n = 1094): metoprolol, ramipril, or amlodipine; no significant difference in GFR slope; ACE inhibitor showed 22% improvement in composite GFR, ESRD, or death versus beta blocker GEMINI (n = 1235): carvedilol versus metoprolol; 16.2% reduction in microalbuminuria and 41% reduction in progression to microalbuminuria	No evidence of preservation of renal function
Angiotensin-converting enzyme (ACE) inhibitor	AASK (n = 1094): metoprolol, ramipril, or amlodipine; no significant difference in GFR slope; ACE inhibitor showed 38% improvement in composite GFR, ESRD, or death versus CCB ALLHAT (n = 33,357): chlorthalidone, amlodipine, or lisinopril versus doxazosin; ACE inhibitor	Reduction in proteinuria Preservation of rena function
	showed no significant difference in the GFR slope during 4 years compared with diuretic (-0.019 versus - 0.018 dL/mg per year) MICRO-HOPE (n = 3577): ramipril versus placebo; 24% reduction in overt nephropathy (> 300 mg proteinuria), with and without baseline microalbuminuria REIN (n = 352): ramipril versus placebo; decline in GFR per month significantly lower with ACE inhibitor versus placebo (0.53 versus 0.88 mL/min), 55% reduction in risk of doubling of baseline	
Angiotensin receptor blockers	creatinine or ESRD	Reduction in proteinuria Preservation of rena
(ARBs)	IDNT (n = 1715): irbesartan or amlodipine versus placebo; 20% reduction in doubling of creatinine, ESRD, creatinine > 6 mg/dL, mortality compared with placebo and 23% compared with CCB MARVAL (n = 332): valsartan versus amlodipine; urine albumin excretion was 56% of baseline with valsartan and 92% of baseline with amlodipine; 29.9% valsartan versus 14.5% amlodipine	function
	reverted to normoalbuminuria RENAL (n = 1513): losartan versus placebo; reduction in time to doubling of creatinine, ESRD, death by 16%, incidence of doubling of serum creatinine by 25%, ESRD by 28%; no impact on mortality TRANSCEND (n = 5927): telmisartan versus placebo; no significant difference in composite dialysis or creatinine doubling or change in GFR	
Combination		
ACE inhibitor/ARB	CALM (n = 199): candesartan, lisinopril, or combination; 50% reduction in urinary ACR with combination compared with candesartan 24% and lisinopril 39% ONTARGET (n = 8576): ramipril or telmisartan versus combination; increased ESRD, creatinine doubling, death in combination 14.5% compared with ACE inhibitor or ARB 13.5%; increase in urinary albumin excretion was 7% less in combination compared with ACE inhibitor VALERIA (n = 133): valsartan/lisinopril versus valsartan or lisinopril; 34% reduction in urine ACR in combination compared with lisinopril; 38% of combination group had normalization of microalbuminuria compared with 17% lisinopril	Reduction in proteinuria Unclear effect on renal function
Direct renin inhibitor (DRI) + ARB	AVOID (n = 599): aliskiren/losartan versus losartan; 20% reduction in urinary ACR without significant reduction in blood pressure	renal function
ACE inhibitor or ARB/ mineralocorticoid receptor blocker (MRB)	Epstein et al (n = 268): eplerenone/enalapril versus enalapril; reduction in urine albumin excretion by 48.4% versus 7.4% Mehdi et al (n = 81): spironolactone/lisinopril versus lisinopril/losartan; reduction in urine albumin excretion with ACE inhibitor/MRB of 34% compared with ACE inhibitor/ARB 16.8%	Reduction in proteinuria Unclear effect on renal function
ACE inhibitor or ARB/ CCB	ASCOT (n = 19,257): perindopril/amlodipine versus atenolol/bendroflumethiazide; 15% reduction in the development of renal impairment Fogari et al (n = 453): fosinopril, amlodipine, or combination; 67% reduction of urine albumin excretion with combination versus fosinopril 46% or amlodipine 33% at 48 months	Reduction in proteinuria Possible effect on renal function





IABLE 11—3 Comparison of Effects of Antihypertens					
Antihypertensive Drug Class	PRC	PRA	Angle I	Angle II	Aldosterone
Diuretic (thiazide)	Т	Т	Т	Т	Т
Beta blocker	44	4 4	4 4	4 4	4 4
Calcium channel blocker	Т	Т	Т	Т	Т
Angiotensin-converting enzyme (ACE) inhibitor	TTT	TTT	TTT	4 4	4
Angiotensin receptor blocker (ARB)	TTT	TTT	TTT	TTT	4 4
ACE inhibitor/ARB	TTT	TTT	TTT	= T	4
Direct renin inhibitor (DRI)	4 4	4 4	4 4	4 4	4
DRI/ARB	4	4	4	4	44
Mineralocorticoid receptor blocker	tut	tut	tut	tut	tut

Ang, angiotensin; PRA, plasma renin activity; PRC, plasma renin concentration.

Renal Effects

Aside from altered lipid profiles and glycemic control, the SHEP and International Nifedipine GITS (INSIGHT) studies showed a greater decline in renal function with diuretic therapy, assessed by a rise in serum creatinine. ^{13,17} The proposed mechanisms of renal injury by diuretics are varied and include volume depletion leading to renal ischemia and stimulation of the RAAS, arteriolar vasoconstriction from endothelial dysfunction and increased oxidative stress, metabolic abnormalities leading to renal hypertrophy and fibrosis, and direct toxic effects on the distal tubular cells ¹⁸ (Table 11-3).

Summary

Diuretics are an acceptable therapy for uncomplicated hypertension. The substitution of hydrochlorothiazide with chlorthalidone in clinical practice may be indicated because of its better tolerability (longer half-life). In younger hypertensive patients and in patients with comorbid conditions, especially diabetes or when the pathophysiology of hypertension may be driven by activation of the RAAS or sympathetic activation, agents other than diuretics or diuretic combinations may be preferred (see Table 11-1).

NITRATES

Nitrates has prodrugs; their active metabolite is nitric oxide, also known as endothelium-derived relaxing factor. Nitrates relax vascular smooth muscle and have vasodilatory effects in both the systemic arteries, including the coronary arteries and collateral vessels, and the veins. ^{19,20} The predominant venodilator effect of nitrates reduces ventricular preload, which in turn reduces myocardial wall stress and oxygen requirements. ²¹ Higher doses of nitrates may also decrease systemic arterial pressure. Other beneficial effects include improved endothelial function and a reduction in platelet adhesion and aggregation. ^{22,23}

Role of Nitrates

Rapid anginal improvement and the resolution of electrocardiographic signs of ischemia have promoted the use of nitrates for the treatment of angina. Trials comparing long-acting nitrates with both calcium channel blockers and beta blockers have shown no significant difference in antianginal efficacy. ²⁴ In the treatment of heart failure, the use of nitrates reduces secondary pulmonary hypertension and improves heart failure symptoms. There are few prognostic data on nitrate therapy in heart failure. The effect of vasodilator therapy on mortality in a chronic congestive heart failure (CHF) trial (V-HeFT I) showed

that when oral nitrates are used with an arterial vasodilator, hydralazine, there is a 34% reduction in overall mortality compared with alpha blockade or placebo. ²⁵ The V-HeFT II trial compared the same combination of nitrate and hydralazine with enalapril in patients with symptomatic CHF. ²⁶ Enalapril showed a further 28% reduction in mortality over combination therapy.

For post-myocardial infarction patients in the fibrinolytic era, trials have failed to show a significant benefit of short term, early nitrate therapy. In the Gruppo Italiano per lo Studio della Sopravvivenza nell'Infarto Miocardico trial (GISSI-3), administration of transdermal nitroglycerin for 6 weeks after acute myocardial infarction demonstrated a non-statistically significant 6% risk reduction in overall mortality. ²⁷ In the Fourth International Study of Infarct Survival (ISIS-4) trial, administration of oral mononitrate for 1 month after acute myocardial infarction also did not demonstrate an impact on postinfarction mortality or combined endpoint of death and heart failure or extensive ventricular dysfunction. ²⁸ The duration of nitrate therapy and follow-up of these trials were relatively short, and the open-label use of nitrates was high in the placebo arms.

With contemporary percutaneous revascularization, one early trial evaluated the long-term nitrate treatment of post myocardial infarction patients and suggested an increase in cardiac events. ²⁹ Nitrate rebound during nitrate-free intervals and tolerance have been suggested as mechanisms to blunt hemodynamic effects and to increase cardiovascular events. There is also the suggestion of reflex neurohormonal activation with an increase in plasma renin activity and circulating angiotensin II (A II) levels during nitrate therapy, which attenuates the vasodilator effect of nitroglycerin during prolonged treatment. 30 Both the rebound phenomenon and tolerance may be suppressed by the use of ACE inhibitor or angiotensin receptor blocker (ARB) therapy. 31-33 More recent retrospective studies using optimal medical therapy have not identified an increase in adverse events, but a more neutral effect, further questioning the utility of nitrates in secondary prevention ^{34,35} (see Table 11-2).

Summary

Nitrates are effective antianginal agents used for the symptomatic therapy of all entities of myocardial ischemia.

therapy can be effective in patients with coronary vasospasm or heart failure caused by myocardial infarction. Their role in the treatment of hypertension is limited. In post-myocardial infarction patients without evidence of myocardial ischemia or heart failure, the role of nitrate therapy in secondary prevention is not well established (see Table 11-2).

CALCIUM CHANNEL BLOCKERS

Calcium channel blockers (CCBs) are classified into the dihydropyridines and the nondihydropyridines, which include phenylalkylamines and benzothiazepines. Examples of the dihydropyridines are nifedipine, felodipine, and amlodipine; examples of the nondihydropyridines are diltiazem and verapamil. Advanced generations of CCBs have modified release formulations, and specifically the newer generation dihydropyridines have higher vascular selectivity and less direct cardiac effects in comparison to short-acting nifedipine.

The CCBs act by counteracting excess calcium entry through the voltage- and receptor-operated calcium channels. CCBs cause potent arteriolar vasodilation by the reduction of calcium entry into the vascular smooth muscle. Within the cardiac myocytes, maintaining the cytosolic calcium balance and preserving ATP stores help maintain myocyte viability and delay ischemic damage. ³⁶ Other effects of CCBs include the protection of endothelial integrity and reduction of endothelial apoptosis, ³⁷ improved nitric oxide production and release, ³⁸ improved fibrinolytic function through an increase in tissue-type plasminogen activator (t-PA) activity, ³⁹ and importantly, reductions in A II-mediated vasoconstriction and decreased A II stimulatory effect on aldosterone production. ⁴⁰ Many of these effects have also been attributed to inhibitors of the RAAS.

Role of Calcium Channel Blockers

Chronic Stable Angina

CCBs dilate the coronary arteries and increase coronary blood flow. Coronary blood flow is one determinant among others, including heart rate, contractility, and arterial pressure, that affects myocardial oxygen consumption, and all can be variably affected by CCBs. For chronic stable angina, three trials have evaluated the long-term effects of CCBs. Angina Prog nosis Study in Stockholm (APSIS) and Total Ischemic Burden European Trial (TIBET) compared verapamil with metoprolol and nifedipine SR with atenolol, respectively, in patients with stable angina pectoris. ^{41,42} Neither study found a difference in morbidity and mortality rates between CCBs and beta blockers. When long-acting nifedipine was added to optimal medical therapy including beta blockade, no difference was noted in mortality, with small reductions in the secondary endpoints of need for coronary revascularization and heart failure. ⁴³

Post-Myocardial Infarction

The Secondary Prevention Reinfarction Israeli Nifedipine trials (SPRINT and SPRINT II) evaluated the effect of the dihydropyridine nifedipine on cardiovascular morbidity and mortality during weeks 2 to 3 for SPRINT and within the first week for SPRINT II after an acute myocardial infarction. No significant difference was noted in morbidity or mortality for late therapy; however, early institution of calcium channel blockade was associated with an increased early mortality (within the first 6 days).

The secondary Danish Verapamil Infarction Trial (DAVIT II) showed for the first time that a nondihydropyridine reduces the incidence of death and reinfarction by approximately 20% when it is started 1 week after acute myocardial infarction. ⁴⁴ A significant reduction in mortality (11.8% versus 7.7%) was noted only in patients without CHF. Interestingly , the order of magnitude of this benefit was similar to that achieved in

numerous beta blocker trials. In the Multi center Diltiazem Post-Infarction Trial (MDPIT), diltiazem nonsignificantly decreased cardiac mortality or nonfatal kidney infarction by 11%. $^{\rm 45}$ However, there was a significant increase in late CHF (21% versus 11%) and an increase in cardiac events (hazard ratio, 1.41) in patients with ejection fractions less than 40%. The diltiazem-associated rise in the frequency of CHF was progressively greater with worsening decrements of ejection fraction.

Congestive Heart Failure

As potent vasodilators, CCBs have been shown to improve hemodynamic responses in CHF patients with notable decreases in pulmonary capillary wedge pressure, systemic vascular resistance, and end-diastolic ventricular pressure. 46 CCBs are also negative inotropic agents and have varying degrees of impairment in cardiac function. The differences in negative 11 inotropy are due to differences in the binding affinity ratios of smooth to cardiac muscle, with dihydropyridines being more vascular selective. An early generation CCB, nifedipine, showed clinical effects not consistent with their hemodynamic benefit, with a significant increase in heart failure deterioration. 47 This has also been noted with the use of diltiazem after myocardial infarction in patients with ventricular dysfunction (ejection fraction < 40%). 48 CHF patients who deteriorated after a CCB have been shown to have a worse prognosis, with a significant decrease in mortality at 1 year. 49

Later generations of CCBs, felodipine and amlodipine, were developed with predominant peripheral vasodilating effects, having minimal effect on the myocardium and with longer half-lives thought to decrease reflex neurohormonal activation in patients with heart failure. The Prospective Ran domized Amlodipine Survival Evaluation trials (PRAISE-1 and PRAISE-2) evaluated the long-term use of amlodipine in advanced heart failure patients. ^{50,51} Patients had New York Heart Association (NYHA) Class III or Class IV heart failure and ejection fractions less than 30% despite treatment with ACE inhibitor, diuretic, and digoxin. PRAISE-1 evaluated 1153 patients regardless of the cause of heart failure, and PRAISE-2 evaluated 1652 nonischemic cardiomyopathy patients. In both trials, amlodipine showed a neutral effect on mortality in patients with severe chronic heart failure.

A trial similar to PRAISE was performed with felodipine, Vasodilator-Heart Failure Trial (V-HeFT III trial). ⁵² This study evaluated 450 patients with NYHA Class II or Class III heart failure and ejection fractions less than 45%. Exercise capacity was not statistically significant, and no differences were noted in mortality or rates of hospitalization (see Table 11-1).

Renal Effects

CCBs have pleiotropic effects that might contribute to renal protection against hypertension-induced damage. These effects include a reduction in mesangial inflammation and proliferation and the promotion of nephrotoxic free radical removal. ⁵³⁻⁵⁵ At the renovascular level, CCBs preferentially dilate the glomerular afferent arteriole, with only modest action on the efferent arteriole. ⁵⁶ This preferential effect, potentially noted more predominantly with dihydropyridine CCBs, may cause glomerular hypertension, increasing capillary intraglomerular pressure that could be associated with progression of renal disease. ⁵⁷⁻⁵⁹

Given their diversity of renal effects and the potential class differences between CCBs on intraglomerular pressures, trials evaluating the effects of CCBs on nephropathy are equally



166 divergent. The African American Study of Kidney Disease and Hypertension (AASK) showed a lesser degree of renal function preservation in African Americans with hypertensive nephrosclerosis treated with amlodipine than with the ACE inhibitor ramipril. ⁶⁰ Furthermore, in this population of patients, amlodipine showed a significant increase in proteinuria compared with both metoprolol and ramipril. The ALLHAT study, on the other hand, showed that amlodipine maintained glomerular filtration rate (GFR) better than therapy with a diuretic or ACE inhibitor. ¹⁵ Overall, the use

I of CCBs in renal disease is safe without any evidence of further deterioration of renal function. CCBs may be better than diuretics and beta blockers, but they are not as effective as RAAS inhibitors in reducing proteinuria and preventing 11 the progression of renal disease (see Table 11-2).

Summary

In patients with chronic stable angina, CCBs may be considered for adjunctive treatment but not as monotherapy unless beta blockers and RAAS inhibitors are not tolerated. Early administration of nifedipine in acute myocardial infarction, unless it is specifically indicated in vasospastic angina, is contraindicated. The nondihydropyridines, possibly because of their heart rate-lowering properties, are the only CCBs to show benefit in the post-myocardial infarction population. They also increase the risk for subsequent CHF in post myocardial infarction patients with reduced ejection fractions and should probably be reserved for patients without heart failure who cannot tolerate a beta blocker or in whom a beta blocker is contraindicated. Amlodipine and felodipine, given their neutral effect on cardiac morbidity and mortality and ease of tolerance, may be considered in the management of hypertension or coronary artery disease (CAD) in patients with heart failure (see Table 11-1).

BETA BLOCKERS

Beta blockade reduces myocardial workload and oxygen demand through a reduction in heart rate, blood pressure, and cardiac index. ⁶¹ A decrease in sympathetic activity reduces fatty acid metabolism and reduces catecholamine levels, helping to redistribute flow to ischemic territories. In addition, beta blockers moderately lower plasma renin activity , even in patients receiving treatment with an ACE inhibitor , in whom the biological effects of renin are already attenuated. ⁶²

Beta blockers differ in the degree to which they antagonize the effects of the sympathetic nervous system. The predominant effect of this class of medications is through interaction with the beta 1 receptor, primarily located within cardiac muscle. However, each drug has varying effects on beta 2 receptor antagonism regulating norepinephrine release, intrinsic sympathomimetic activity (which provides a mea surable beta agonist response), and other adrenergic receptors such as the alpha 1 receptor. Nonselective beta antagonists block beta 1 and beta 2 receptors; an example is propranolol. Selective beta antagonists block beta 1 receptors; examples include atenolol metoprolol, bisoprolol, and nebivolol. Carvedilol and nebivolol block three adrenergic receptors (alpha 1, beta 1, and beta 2) and reduce endothelin and oxidative stress. 63

Role of Beta Blockers

For decades, beta blockers have been widely used in the treatment of uncomplicated hypertension. However, no trial has shown that lowering blood pressure with a beta blocker reduces the risk of cardiovascular events in patients with hypertension compared with placebo. More recently, beta blocker treatment has been associated with a substantially higher (16%) risk of stroke than treatment with other antihypertensive agents. ⁶⁴ A

proposed mechanism may be differences in the hemodynamic effects of beta blockade.

Treatment with a beta blocker results in reduced brachial blood pressure but does not lower central aortic blood pressure as much as treatment with an ACE inhibitor, diuretic, or CCB does. ⁶⁵ Central aortic blood pressure may be more predictive of cardiovascular events than traditional brachial blood pressure measurements. ⁶⁶ The outcome of both the Anglo-Scandinavian Cardiac Outcomes Trial-Blood Pressure Lowering Arm (ASCOT-BPLA) and the Losartan Intervention For Endpoint reduction in hypertension study (LIFE) showed a stroke reduction of approximately 25% when treatment with either a CCB or an ARB, respectively, was given instead of a beta blocker. ^{67,68}

Post-Myocardial Infarction and Chronic Stable Angina

Early beta blockade relieves chest pain and reduces infarct size and fatal arrhythmias; long-term beta blockade reduces late mortality and reinfarction. ⁶⁹ Early trials and meta analyzes performed before reperfusion therapy suggested that beta blockade after myocardial infarction can reduce mortality , with the most marked reduction of about 25% occurring in the first 2 days after infarction. ⁷⁰ In the Cooperative Cardiovascular Project, beta blockade showed a mortality reduction of 40% across all patient subgroups, including high-risk groups such as the elderly and patients with a severely reduced ejection fraction.

The two largest pre-thrombolytic era trials examining beta blockade in acute myocardial infarction were the Metoprolol in Acute Myocardial Infarction (MIAMI) and the First International Study of Infarct Survival (ISIS-1) trials. ^{72,73} In the MIAMI trial, more than 5000 post-myocardial infarction patients received metoprolol within 24 hours of presentation. Mortality was 4.9% in the placebo group and 4.3% in the metoprolol group. Despite the overall nonsignificant reduction in mortality, subgroup analysis revealed a survival advantage of nearly 30% in patients considered at higher risk. ISIS-1 trial evaluated the efficacy of early atenolol in nearly 16,000 patients presenting with myocardial infarction within 12 hours of symptoms. This trial showed a 15% reduction in vascular deaths as well as a 15% reduction in early cardiac arrests and reinfarctions, predominantly noted within the first day after infarction.

This benefit of beta blockade was evident in early administration and short-term continuation, before the standard use of fibrinolytic and antiplatelet therapy. It can be attributed to the prevention of life-threatening arrhythmias and cardiac rupture. The ClOpidogrel and Metoprolol in Myocardial Infarction Trial (COMMIT) is the largest trial to date, including nearly 46,000 patients to evaluate the effects of intravenous and then oral metoprolol in acute myocardial infarction added to current antiplatelet and fibrinolytic therapy. 74 There was no significant difference noted in the rate of composite endpoint of death, reinfarction, or cardiac arrest, with overall mortality balanced by a 22% reduction in late arrhythmic death and a 29% proportional increase in early shock-related death. The overall net effect of metoprolol therapy changed from being significantly adverse during the first 2 days after admission due to shockrelated death to being significantly beneficial thereafter due to a reduction in ventricular fibrillation.

In combining a retrospectively defined low-risk subgroup of COMMIT similar to the populations studied in MIAMI and ISIS-1, the overall effects were consistent, showing a significant reduction of 13% in mortality, 22% in reinfarction, or 15% in ventricular fibrillation. Beta blocker therapy for these trials among others, including the Beta Heart Attack Trial

(BHAT) and the TIMI IIB trial, was continued up to 2 years. 75.76 Furthermore, randomized trials of beta blocker therapy in patients undergoing percutaneous coronary intervention without fibrinolytic therapy have not been performed. It does seem reasonable, however, to extrapolate the results from those receiving another form of revascularization to the per cutaneous coronary intervention population.

The use of beta blockade for the treatment of chronic stable angina has also been predominantly extrapolated from the evidence of improved mortality of beta blockers in the post myocardial infarction patient. Given the lack of mortality benefit and increased risk of stroke in patients with uncomplicated hypertension, treatment of patients with chronic stable angina but no prior myocardial infarction with beta blockers is not well supported but commonly used. Other antihypertensive medications, such as CCBs and ACE inhibitors, may be equally beneficial without the potential adverse effects of beta blockade therapy.

Congestive Heart Failure

Because of transient negative inotropic effects of beta block ade, beta blockers used to be contraindicated in CHF. Many trials have since shown that patients with heart failure derive greater benefit from beta blocker therapy after myocardial infarction than do patients without heart failure. In the BHAT trial, propranolol reduced mortality equally in patients with and without systolic dysfunction by approximately 25%. However, there was a 47% reduction in sudden death in patients with heart failure, compared with a 13% reduction in patients without heart failure.

The Metoprolol CR/XL Randomized Intervention Trial in Congestive Heart Failure (MERIT-HF), Cardiac Insufficiency Bisoprolol Study II (CIBIS II), and Carvedilol on Survival in Severe Chronic Heart Failure (COPERNICUS) trials evaluated the effects of beta blockade in patients with heart failure, with severely reduced ejection fractions, and receiving stable ACE inhibitor and diuretic therapy. 77-79 Metoprolol, bisoprolol, and carvedilol, respectively, showed an overall reduction in total mortality of approximately 35%. In addition to improved mortality, beta blockers, when given with ACE inhibitors, improve patient symptoms and NYHA functional class and reduce heart failure hospitalizations (see Table 11-4).

Renal Effects

Beta $_1$ receptors alter the release of renin, and beta $_2$ and alpha $_1$ receptors alter sympathetic activity responsible for renovas cular tone, important in the genesis of hypertension and progression of kidney disease. Although beta blockers improve cardiac mortality, they have not shown the same benefit on renal outcomes compared with other agents that lower blood pressure and albuminuria. Studies in both diabetic and non-diabetic nephropathy do not show a significant reduction in microalbuminuria with beta blockade compared with ACE inhibition. 80,81

In the Glycemic Effects in Diabetes Mellitus: Carvedilol-Metoprolol Comparison in Hypertensives (GEMINI) trial, 1235 hypertensive, diabetic patients were enrolled to evaluate the effects of metoprolol versus carvedilol on glycemic control, with a secondary endpoint of change in microalbuminuria. 82 There was a 16.2% reduction in microalbuminuria for patients randomized to carvedilol compared with meto prolol despite similar blood pressure control during 3 years. Among patients with normal albuminuria at baseline, fewer in the carvedilol group (6.6%) than in the metoprolol group (11.1%) progressed to microalbuminuria. GEMINI suggests that beta blockers with alpha 1 receptor activity may reduce the compensatory stimulation of the sympathetic system and RAAS caused by beta blockade and may further reduce the

TABLE 11—4 Compelling Indications for Antihypertensive Drug Classes

Compelling Indication	Therapeutic Options
High CAD risk	Thiazide*, beta blocker, CCB*, ACE inhibitor*, ACE inhibitor/CCB, ARB/ CCB
Chronic stable angina	Nitrates, CCB, ACE inhibitor, ACE inhibitor/CCB, ARB/CCB
Post-myocardial infarction	Beta blocker*, ACE inhibitor*, ARB, MRB*, nondihydropyridine CCB
Heart failure	Beta blocker*, ACE inhibitor*, ARB*, MRB*, diuretics, nitrates

Chronic kidney disease

Reduction in proteinuria Beta blocker, ACE inhibitor*, ARB*,

DRI, ACE inhibitor/ARB, ACE

inhibitor/MRB, ARB/MRB, ACE inhibitor/CCB, ARB/CCB Preservation of renal CCB, ACE inhibitor*, ARB*, ACE function inhibitor/CCB, ARB/CCB

*Classes recommended by the Seventh Report of the Joint National Committee (JNC7).
ACE, angiotensin-converting enzyme; ARB, angiotensin receptor blocker; CAD, coronary artery disease; CCB, calcium channel blocker; DRI, direct renin inhibitor; MRB, mineralocorticoid receptor blocker.

progression of albuminuria, providing cardiorenal protection (see Table 11-2).

Summary

Beta blockers have an established benefit for the treatment of patients after myocardial infarction or with heart failure and play an important role in secondary prevention. The role of beta blockade in the treatment of uncomplicated hypertension remains unclear, given the paucity of data and the possible mild increased risk of stroke. Delay of the initiation of beta blocker therapy until after stabilization of the patient from an acute myocardial infarction may avoid the excess risk of shock and early mortality while preserving much of its described benefits. Controversy remains as to the duration of beta blocker therapy in post-myocardial infarction patients, but if the medication remains well tolerated, discontinuation should be discouraged (see Table 11-4).

RENIN-ANGIOTENSIN-ALDOSTERONE Syst

Renin is a protease produced by the juxtaglomerular cells of the kidney that line the afferent arteriole of the glomerulus. Renin is secreted mainly as a result of four mechanisms: renal baroreceptors in the afferent arteriole that sense alterations in renal perfusion pressure, a change in NaCl delivery to the macula densa cells of the distal tubule, sympathetic stimulation through beta 1-adrenergic receptors, and negative feedback by A II on juxtaglomerular cells. ⁸³ Outside of the kidney, renin is secreted by other tissues including the brain, adrenal gland, ovary, and adipose tissue in addition to the heart and vascular tissue. Renin is the rate-limiting step of the RAAS (Fig. 11-1).

Renin cleaves angiotensinogen to angiotensin I (AI), commonly referred to as Ang-(1-10) for its decapeptide structure. Angiotensinogen is primarily produced by the liver, but expression has been noted in tissue similar to that expressing renin, including the kidney and heart. AI is biologically inactive and is converted by ACE by the removal of a dipeptide to





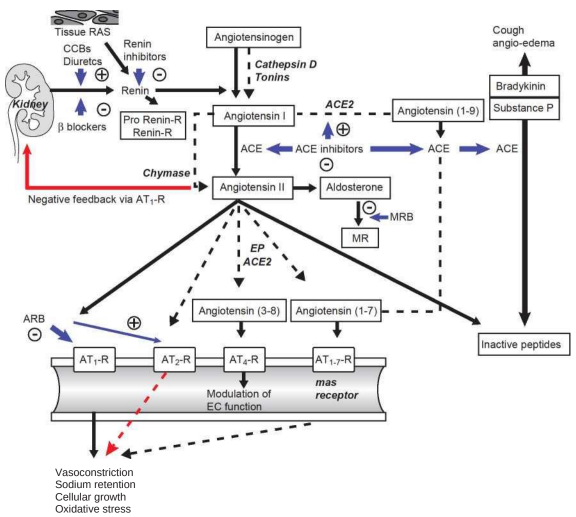


FIGURE 11-1 The renin-angiotensin-aldosterone system. ACE, angiotensin-converting enzyme; ARB, angiotensin receptor blocker; AT-R, angiotensin receptor; CCBs, calcium channel blockers; EC, endothelial cells; EP, endopeptidases; MR, mineralocorticoid receptor; MRB, mineralocorticoid receptor blocker. Black arrows show stimulation, red arrows show inhibition, and blue arrows show antihypertensive drug involvement with stimulation (+) or inhibition (-) of the pathway. Full lines represent major functional pathways, and dotted lines show alternative pathways. (Modified from Staessen J, Li Y, Richart T: Oral renin inhibitors. Lancet 368:1449, 2006.)

form A II or Ang-(1-8). ACE is a membrane-bound exopeptide predominantly localized on the membranes of the vascular endothelial and renal proximal tubular cells. ACE also metabolizes other vasodilator peptides, including bradykinin and kallidin, which may have important physiological effects that will be described later in the chapter.

Although A II is the primary active product in the RAAS, other metabolites of AI and A II may have physiological activity . These include angiotensin III, (Ang-[2-8]); angiotensin IV, (Ang-[3-8]); and Ang-(1-7). All metabolites are formed by removal of varying amounts of amino acids from either the N or C terminus of A II and have varying effects, some of which are unknown to

It is the interaction of A II with the angiotensin receptors that drives many of the RAAS effects. 83 At least four angiotensin receptors have been described; the AT 1 receptor mediates many of the pathophysiological effects and is therefore a major therapeutic target for secondary prevention. AT 1 receptors are expressed in the kidney, adrenals, liver, brain, vasculature, and heart, and their activity includes vasoconstriction, vascular and cardiac hypertrophy, renal tubular sodium resorption, inhibition of renin release, sympathetic activation, cell growth and proliferation, and increased inflammatory response. AT 2 receptors are not highly expressed in the adult and may oppose some of the AT 1 receptor-mediated pathways. Some physiological effects of AT 2 recep tors may include vasodilation, smooth muscle antiproliferation, and inhibition of cardiac remodeling. AT 3 receptor function is unknown, and AT 4 receptor function has been shown to modulate endothelial function through plasminogen activator inhibitor 1 (PAI-1). 83

A II also stimulates, through the AT 1 receptor, the release of aldosterone, predominantly from the adrenal cortex, but extraadrenal sites of aldosterone synthesis have been identified in the brain, vasculature, and heart. Aldosterone is regulated by both A II and dietary potassium. 84 It is an important hormone in sodium and fluid retention as a result of direct stimulation of the mineralocorticoid receptors in the distal renal tubules. Mineralocorticoid receptors are also as wide spread as their extra-adrenal sites. Their varied proposed physiological mechanisms include renal, vascular, and cardiac inflammation; fibrosis and hypertrophy; increases in sympathetic activation; attenuation of platelet aggregation; and impairment of endothelial dysfunction 85,86 (Fig. 11-2).

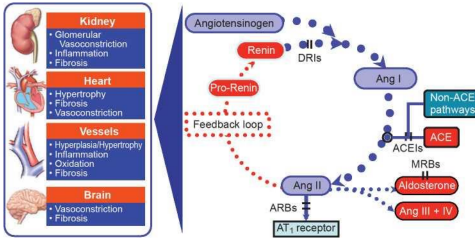


FIGURE 11-2 Classic understanding of the renin-angiotensin-aldosterone system with feedback loop activation. ACE, angiotensin-converting enzyme; ACEIs, angiotensin-converting enzyme inhibitors; Ang, angiotensin; ARBs, angiotensin receptor blockers; AT,, type 1 angiotensin II receptor; DRIs, direct renin inhibitors; MRBs, mineralocorticoid receptor blockers.

Intracellular Renin-Angiotensin-Aldosterone System

Despite the multiorgan involvement, specifically in the kidney, vasculature, and heart, for the full physiological axis of the RAAS, local tissue or intracellular RAAS has been identified and is believed to play a significant role in cardio vascular pathophysiology. ⁸⁷ The intracellular RAAS is defined by a functionally active complete RAAS within a cell, including components that participate in A II production and its receptors and effector proteins. In the heart, angiotensinogen is internalized and basal expression of RAAS components is very low. Intracellular RAAS is activated in response to pathological conditions such as pressure or volume overload , myocardial infarction, and diabetes. ⁸⁸

Potential mechanisms of action include direct intracellu lar A II effects on the heart, which are non-AT $_{\rm I}$ receptor mediated, and intracellular A II regulation of the AT $_{\rm I}$ receptor , providing an improved response to extracellular A II. These mechanisms have been suggested to have a role in left ventricular hypertrophy, activation of metalloproteinases with facilitation of plaque rupture, and pathogenesis of vascular disease. 88 Intracellular A II actions are not blocked by ACE inhibitors or ARBs because of their inability to enter the cell and to block these pathways. More complete RAAS inhibition may be provided by direct renin inhibitors that can block both extracellular and intracellular pathways.

Alternative Pathways of Angiotensin I Production

In addition to intracellular RAAS, alternative pathways of A II production independent of ACE, termed ACE escape, have been established. These include chymase and ACE-2-dependent pathways. Chymase is a protease stored within the secretory granules of mast cells; it is present within the blood vessels, the heart, and other tissues but does not circulate within the blood. In blood vessels, chymase is localized to mast cells in the adventitia, whereas ACE is localized in the vascular endothelium. ⁸⁹ In normal vasculature, circulating AI does not readily penetrate into the interstitial space, and hence AI is converted to A II in an ACE-dependent fashion. In the setting of vascular injury or inflammation, mast cells are activated, allowing a chymase-dependent conversion of AI to A II.

Although chymase may only have a limited physiological role in the generation of A II, residual A II formation may be observed in the presence of ACE inhibitor due to this chymase-dependent pathway. Other effects of chymase, outside of the A

II-mediated effects, include promotion of atherosclerosis through the prevention of cholesterol removal by HDL, intimal hyperplasia of injured arteries, potential erosion and rupture of coronary atheroma leading to an acute coronary syndrome, and stimulation of transforming growth factor- $\mbox{\cite{S}}$ (TGF- $\mbox{\cite{S}}$) leading to cardiac fibrosis. 90

Angiotensin-converting enzyme 2 (ACE-2) is a peptidase present throughout the cardiovascular system, including in macrophages, vascular endothelium, smooth muscle cells, and potentially cardiac myocytes. It has been shown to convert AI and A II to Ang-(1-9) and Ang-(1-7), respectively . ⁹¹ The full physiological importance of this pathway has yet to be determined. However, ACE-2 may have a counter-regulatory role with ACE in fine-tuning the RAAS system through increased production of Ang-(1-7), balancing A II vasopressor effects. ⁹²

Kallikrein-Kinin System

The main effector protein of the kallikrein-kinin system is bradykinin, produced by the proteolytic cleavage of a highmolecular-weight kininogen by kallikrein. Bradykinin has a variety of biological effects, some cardioprotective. These effects are mediated through BK 1 and BK 2 receptors located on endothelial cells. 93 The BK 1 receptor is rarely expressed in normal tissue but is upregulated in pathological states associated with inflammation and tissue injury. 94 It causes stimulation of smooth muscle cells, cell proliferation, and collagen synthesis. The BK 2 receptor, on the other hand, leads to increased stimulation of nitric oxide and prostacyclin, exerting vasodilator, anti-ischemic, and antiproliferative effects. 95 Kinins are released in the setting of ischemia, myocardial infarction, and heart failure, and the ability of ACE inhibitor to prevent the breakdown of kinins represents a potential mechanism contributing to the drug's cardioprotective effects. 96,97

Prorenin and Prorenin Receptor

Renin is synthesized as a prohormone, prorenin, and stored in granules within the renal juxtaglomerular cells. ⁹⁸ Prorenin is activated by removal of its N terminus that covers the active



FIGURE 11-3 Interaction of cardiovascular and renovascular disease. DM, diabetes mellitus; HTN, hypertension.

site and blocks access to angiotensinogen. ⁹⁹The recent discovery of the prorenin receptor has expanded the physiological role of this once believed inactive prohormone. ¹⁰⁰The prorenin receptor binds to both renin and prorenin and has been localized to the vascular smooth muscle, the heart, and the distal and collecting tubules of the kidney.

When renin binds to the receptor, its activity is amplified fivefold, accelerating the production of AI and in turn A II. Prorenin, when it is bound to the receptor, can undergo non-proteolytic activation by unfolding of the peptide, exposing its active site. This allows prorenin to assume full activity and to contribute to the production of A I. Independent of A II, both renin and prorenin exert physiological effects through the prorenin receptor. ¹⁰¹ The activated receptor stimulates the production of TGF- \$ and heat shock proteins, increasing levels of PAI-1 and collagen involved in organ fibrosis and actin filament dynamics involved in contractility, hypertrophy, and apoptosis. ¹⁰²

Plasma Renin Activity

Renin is the primary effector hormone of the RAAS, the first and rate-limiting step in A II synthesis. Plasma renin concentration is a measure of the amount of renin in the circulation, and plasma renin activity is a measure of its catalytic activity. 98 Plasma renin activity correlates with the rate at which AI is generated and is a marker of the activation of the RAAS.

Early data have identified that elevated plasma renin activity has a direct relationship with adverse cardiovascular outcomes, including CHF and death, particularly in patients with CAD. ^{103,104} More recently, several studies by the Inter mountain Heart Collaborative Study Investigators have demonstrated a correlation of plasma renin activity to both worsened ischemia and heart failure outcomes at 3 years in patients with mild and moderate coronary atherosclerosis. ^{105,106} The use of ACE inhibitors or ARBs can increase plasma renin activity by inhibiting the feedback loop that governs the activity of the RAAS. The role and importance of prorenin, plasma renin concentration, and plasma renin activity have raised interest in the use of direct renin inhibitors (see Table 11-3).

Association Between Renal Insufficiency and Cardiovascular Disease

Many studies have demonstrated that the relationship between renal dysfunction and increased cardiovascular morbidity and mortality extends across the spectrum of renal dysfunction to include even the mildest degree of renal impairment. ¹⁰⁷ This relationship is also maintained across varying degrees of baseline cardiovascular health. In the general population ¹⁰⁷⁻¹⁰⁹ and in patients with hypertension, ¹¹⁰ stable CAD, ^{111,112} postmyocardial infarction, ¹¹³ or heart failure, ¹¹⁴ there is an independent and strong correlation between worsening renal function and an increase in cardio vascular events. Baseline GFR has even been suggested to be a stronger predictor of mortality

than either left ventricular ejection fraction or New York Heart Association class. 114

The increased cardiovascular risk for individuals with renal insufficiency is explained in part by the increased prevalence and severity of diabetes and hypertension (Fig. 11-3). However, other mechanisms have been proposed to contribute to the increased risk of cardiovascular events in renal insufficiency. These mechanisms include an increase in endothelial dysfunction, an increase in oxidant stress and inflammation, progressive dyslipidemia, and stimulation of the RAAS that accompanies this disorder. ¹¹⁵

In renal disease, the initial insult to the kidney results in a decline in the GFR, noted by an increase in serum creatinine concentration. 116 When nephrons are damaged, the remaining nephrons undergo hypertrophy with alterations in arteriolar tone. This in turn increases hydraulic pressure within the glomerular capillaries. The filtration capacity of the remaining nephrons increases, minimizing the functional consequences of nephron loss. The high glomerular capillary pressure enlarges the glomerular membrane pores through a process partially attributed to angiotensin (A II), which subsequently increases the number of proteins, both albumin and nonselective proteins, filtered. It is these proteins that become reabsorbed by the proximal tubular cells and may stimulate a vasoactive and inflammatory framework leading to a systemic vascular disease that can affect the kidneys as well as other arterial systems in the body, including the heart. Endothelial dysfunction, which is associated with alterations in vasomotor tone, inflammation, and oxidative stress, plays a key role in remodeling of the vasculature and increases the risk for atherosclerosis.

Described mediators of inflammation and coagulation known to be increased during renal insufficiency include C-reactive protein, fibrinogen, homocysteine, interleukin-6, factor VII/VIII, von Willebrand factor, D-dimer, and plasmin-antiplasmin complex. ^{117,118} These mediators, derived in part from increased A II and the modifications of lipids, directly affect endothelial function and vascular remodeling through enhanced oxidative stress. Increased oxidative stress produces reactive oxygen species mainly through the activation of nicotinamide adenine dinucleotide (NADPH) oxidase and the uncoupling of endothelial synthase. The production of reactive oxygen species directly reduces nitric oxide activity, promoting endothelial dysfunction and atherosclerosis.

Dyslipidemia associated with renal insufficiency further contributes to the production of reactive oxygen species and inflammation. Notable alterations in lipid synthesis include elevated levels of lipoprotein(a) and triglycerides, decreased levels of apo A and HDL, and the decreased ability to reduce oxidized, small dense LDL. ¹¹⁷ These defects are the result of a combination of overproduction and clearance defects in apo B-containing lipoproteins. The dyslipidemia of renal disease

is complex, with an atherogenic profile similar to that of metabolic in cardio vascular events and a 27% reduction in the risk of heart syndrome. ¹¹⁹ In general, the severity of dyslipidemia correlates with failure. the severity of proteinuria. 120

highly correlated with the severity of glomerulosclerosis and of hypertension (LIFE), reconfirmed the association between varying renal arteriolosclerosis. 121 Renal insufficiency may be a marker for degrees of proteinuria and cardiovascular risk, independent of generalized atherosclerotic burden. 122 However, the presence of blood pressure. 130,134 Reductions in proteinuria over time translate these altered pathways has been noted in people with renal insufficiency but with no evidence of clinical or subclinical be more predictive than the other traditional cardiovascular risk cardiovascular disease. The prevalence of endothelial dysfunction, factors. Proteinuria can be used as a potential target or indicator of low-grade inflammation, dyslipidemia, and RAAS stimulation therapeutic response with antihypertensive therapy. associated with renal disease may explain the acceleration of atherosclerosis. 117 In combination with hypertension and associated left ventricular hypertrophy, this may lead to a reduction in coronary reserve, explaining the prevalence of coronary ischemia ENZYME INHIBITORS and cardio vascular events in people with renal insufficiency. 118

Association Between **Proteinuria** Cardiovascular Disease

decline in the GFR, an increased rate of albumin or protein excretion left ventricular dilation in combination with a reduction in reflects a derangement in the glomerular filtration barrier. 123 The preload and afterload is beneficial in CAD patients, especially in presence of abnormal quantities of albumin in the urine often the post-myocardial infarction state. ACE inhibitor therapy precedes renal functional deterioration. Microalbuminuria, an alters cardiac remodeling and has a multitude of vascular albumin-to-creatinine ratio (ACR) of 30 to 300 mg/g, correlates with protective effects through decreased levels of A II and nephrosclerosis; macroalbumin uria (ACR > 300 mg/g) generally aldosterone and upregulation of bradykinin. In addition, indicates established renal parenchymal damage.

Proteinuria (or albuminuria) has been shown to be an independent marker of cardiovascular risk. 124-126 The link between balance with reduced PAI-1 and increased t-PA activity have albuminuria and adverse cardiovascular events was first recognized been shown. 136-138 It is the balance of cardiac and vascular with macroalbuminuria, later with microalbuminuria, and currently protective effects that provides the benefit of ACE inhibitor with any evidence of albuminuria. 127-129 It has been suggested that therapy. proteinuria is a continuous cardiovascular risk factor, whereas microalbuminuria is a designated threshold for renal functional deterioration in individuals with and without diabetes. 117,130

Proteinuria, despite its association with renal injury, may also reflect a systemic increase in endothelial permeability or Post-Myocardial Infarction and Congestive Heart Failure dysfunction. 131 The vascular endothelium has an important role in regulating the transport of proteins across vessel walls. Endothelial dysfunction, as a result of shear stress, may allow the deposition of lipoproteins in the subendothelial space, promoting atherogenesis. Further injury to the endothelium results in increased cell and Further injury to the endothelium results in increased cell and with or without symptoms, or the presence of an earlier platelet adhesiveness, greater permeability to inflammatory cells, myocardial infarction. 139-141 The results of these trials were and altered production of vasoactive mediators, specifically nitric oxide. 131 This suggests that proteinuria may be a marker of generalized damage to the peripheral vasculature, paralleling the degree of cardiac organ damage.

The Irbesartan Diabetic Nephropathy Trial (IDNT) and Reduction of Endpoints in NIDDM with the Angiotensin II Antagonist Losartan (RENAAL) trial were the first trials to help establish proteinuria as a cardiovascular risk factor. 132,133 IDNT enrolled subjects with type 2 diabetes, hypertension, and macroalbuminuria. A total of 1715 subjects were randomized to irbesartan, amlodipine, or placebo and were observed for a mean period of 2.6 years. The primary outcome was a composite of serum with captopril in the TRACE study was a significantly reduced creatinine doubling, end-stage renal disease (ESRD), or death. Although irbesartan proved to have a significant benefit compared with other therapies, no difference was detected on a secondary Study (CONSENSUS II), GISSI-3, and ISIS-4 trials further outcome of car diovascular events. Further analysis of baseline albuminuria and cardiovascular composite endpoint showed that 172 evaluated this short-term use of early ACE inhibitor therapy the proportion of patients who experienced cardiovascular events progressively increased with increasing quartiles of albuminuria, a 1.3-fold increased risk for each natural log unit increase in ACR.

RENAAL enrolled the same population of subjects as IDNT 171 did, with a total of 1513 subjects randomized to either losar tan or placebo. Baseline albuminuria was again shown to be a predictor of composite cardiovascular endpoint as well as heart failure alone. In addition, the change in urinary albumin from baseline to 6 months was the only correlate of adverse cardiovascular outcomes; a 50% reduction in base line albuminuria translated into an 18% reduction

Other trials, such as the Heart Outcomes Prevention Evaluation It has been demonstrated that the severity of atherosclerosis is (HOPE) and the Losartan Intervention For Endpoint reduction in into improved outcomes, and it appears that proteinuria may even

ANGIOTENSIN-CONVERTING

After an acute myocardial infarction, a complex of changes and involving neurohormonal activation and cardiac remodeling can lead to progressive heart failure and death. Left ventricular dilation is an integral part of this remodeling process, Whereas an elevated serum creatinine concentration reflects a compensating for the loss of systolic function. 135 Attenuation of modulation of adrenergic tone through the effects on A II, improved endothelial function, and improved fibrinolytic

Role of Angiotensin-Converting Enzyme Inhibitors

Early long-term trials, such as Survival and Ventricular Enlargement (SAVE), Acute Infarction Ramipril Efficacy (AIRE) study, and Trandolapril Cardiac Evaluation (TRACE), enrolled high-risk patients after myocardial infarction with heart failure, consistent and demonstrated that the use of an ACE inhibitor significantly reduced overall heart failure and mortality by about 5% for each endpoint, with a relative reduction of approximately 20%.

In the Survival of Myocardial Infarction Long-term Evaluation (SMILE) trial, a shorter term treatment of ACE inhibitor for 6 weeks was instituted within 24 hours after an acute anterior myocardial infarction. 142 It showed a significant reduction in a combined endpoint of death and severe heart failure by 34%, attributable mainly to a decrease in the incidence of CHF. However, one of the unexpected benefits from therapy likelihood of a subsequent myocardial infarction.

The Cooperative New Scandinavian Enalapril Survival

but expanded the population to include all acute myocardial infarction patients. 143-145 When therapy was instituted within 24 hours, there was a less than 1% improvement in heart failure and mortality, with only GISSI-3 and ISIS-4 showing a statistically significant shortterm survival benefit. Sub group analysis showed, as in the early trials, a more pronounced benefit in patients with left ventricular dysfunction and previous infarctions.

Coronary Artery Disease

ACE inhibitors have been shown to reduce cardiovascular

failure. Unexpected findings from the SAVE and Studies of 11 Left high-risk patients without left ventricular dysfunction. Not all of in patients with chronic heart failure 152 (see Table 11-4). the benefit could be attributed to a reduction in blood pressure because a majority of the patients did not have hypertension at Renal Effects baseline and the mean reduction in blood pressure with treatment was relatively small.

In the European Trial on Reduction of Cardiac Events with Perindopril in Stable CAD (EUROPA) study, patients with lower risk asymptomatic stable CAD (ie, prior myocardial one coronary artery, or an abnormal stress test, without heart failure or substantial hypertension) underwent treatment with pressure and the level of proteinuria. 153 an ACE inhibitor, perindopril, which was shown to reduce cardiovascular events by 20%. 148 This endpoint was predominantly driven by a significant reduction in nonfatal myocardial infarction with a lower rate of heart failure. EUROPA extended the observations of the HOPE study to a population with an improved prognosis, showing an overall similar reduction in cardiovascular events. Last, the nephropathy with ACE inhibitor versus control. Prevention of Events with ACE Inhibition trial (PEACE) evaluated the efficacy of the ACE inhibitor trandolapril for a reduction in atherosclerotic complications among patients ventricular function. 149 There was no significant difference in the primary endpoint of cardiovascular death, myocardial infarction, or coronary revascularization with a median follow-up of almost 5 years.

context of HOPE and EUROPA is important. It has been argued that the population of patients from PEACE underwent more intensive management of baseline risk factors, with a larger proportion of patients treated with lipidaddition, a larger proportion of patients underwent coronary revascularization before enrollment. This can be noted in a minimal adverse events in both diabetic much lower overall event rate in the PEACE trial compared with both EUROPA and HOPE. However, further subgroup analyzes of HOPE and EUROPA reevaluating the endpoints on lipid-lowering agents, beta blockade, and antiplatelet therapy or without prior revascularization showed similar benefits. This analysis suggested that the neutral results of PEACE were probably due not to the lower risk of patients or background therapies but rather to inadequate power of the study and perhaps the relatively low dose of trandolapril that was used.

Timing and Choice of ACE Inhibition

The optimal timing of ACE inhibition after acute myocardial infarction remains controversial. ACE inhibition started relatively late (> 3 days) in high-risk patients showed an important long-term benefit on heart failure and mortality. On the other hand, early institution (< 24 to 36 hours) showed a relatively small short-term benefit, mainly during the first 7 days when mortality was higher. ¹⁵⁰ The cause of death was mainly due to a higher rate of cardiac rupture, electromechanical dissociation, and pump failure after myocardial infarction in the untreated patient. There was no overall effect on reinfarction rates in the timing of therapy, but rates of

events in patients with acute myocardial infarction and heart hypotension were 10% higher with early ACE inhibitor institution.

The choice of ACE inhibitor varies between many of the clinical Ventricular Dysfunction (SOLVD) trials showed a significant and trials. A comparison of enalapril and captopril showed that there consistent reduction in reinfarction rates during long-term follow- was a significant and equivalent benefit between the ACE inhibitors. up. 139,146 Three large clinical trials have further investigated the This trial, among others that had similar populations of patients but impact of ACE inhibition in patients with stable CAD without any different choice of ACE inhibitors, suggested an overall class effect evidence of heart failure or left ventricular dysfunction: HOPE, of ACE inhibition in excess of the benefits of optimal medical EUROPA, and PEACE. In the HOPE study, high-risk patients with therapy. In addition, higher doses of ACE inhibition provide greater vascular disease including CAD or diabetes plus at least one other haemodynamic and symptomatic benefit than low doses do. 151 The cardiovascular risk factor without heart failure were included. 147 Assessment of Treatment with Lisinopril and Survival (ATLAS) trial The primary endpoint of cardiovascular death, myocardial evaluated the effects of low-dose compared with high-dose ACE infarction, or stroke occurred less frequently, approximately 4%, inhibitor and showed a nonsignificant 8% reduction in mortality and with a 22% risk reduction. This trial suggested that ACE inhibitor a significant 12% reduction in mortality or hospitalization. Overall, therapy is effective in the prevention of cardiovascular events in high doses of ACE inhibitor reduce the risk of major clinical events

Much of the antiproteinuric effects of RAAS blockade can be attributed to changes in glomerular hemodynamics. Both ACE inhibitors and ARBs, discussed later, induce preferential vasodilation of the efferent arteriole, decreasing glomerular infarction > 3 months, percutaneous coronary intervention or capillary pressure. Blockade of A II with these agents also coronary artery bypass grafting, > 70% narrowing of at least significantly reduces the inflammatory, proliferative, and fibrotic changes that occur in renal disease, further reducing glomerular

In the Microalbuminuria, Cardiovascular, and Renal Outcomes-Heart Outcomes Prevention Evaluation (MICRO HOPE) trial, nearly 3600 patients with diabetes at high risk for cardiovascular events were treated with ramipril or placebo. 154 In patients without baseline microalbuminuria, the risk of new microalbuminuria was nonsignificantly reduced. There was a 24% reduction in overt

In nondiabetic patients with chronic nephropathy, the Ramipril Efficacy In Nephropathy (REIN) trial also noted a similar benefit. ^{155,156} Patients were randomized to ramipril or conventional therapy with low-risk stable CAD with normal or slightly reduced left with the aim of achieving comparable blood pressure control. Results showed that in patients who had rapid progression of renal disease and macroalbu minuria at baseline, ACE inhibitor safely lowered the rate of decline in GFR and reduced by half the combined risk of creatinine doubling or progression to ESRD. The AASK trial Interpretation of the negative findings of PEACE in the showed a greater preservation of renal function in African Americans with hypertensive nephrosclerosis treated with ramipril than with CCB but no significant improvement in proteinuria. 60

Not all studies with ACE inhibitors demonstrate an improvement in renal function over time. In the ALLHAT study, lowering agents, beta blockade, and antiplatelet therapy. In there was no difference in GFR with an ACE inhibitor or diuretic therapy. ¹⁵ Overall, ACE inhibitors slow renal function decline with and nondiabetic nephropathy with various degrees of albuminuria VALIANT showed that death, reinfarction, or recurrent heart (see Table 11-2).

Summary

The greatest benefit of ACE inhibition is noted in the selective treatment of high-risk patients, in particular patients with acute myocardial infarctions (specifically a large anterior wall), failed The first trial to directly compare an ARB with an ACE inhibitor in reperfusion therapy, and left ventricular dysfunction . ACE inhibition should be considered for all patients with left ventricular dysfunction and CAD and for patients with multiple risk factors including diabetes, metabolic syndrome, and hypertension. The magnitude of benefit from an ACE inhibitor is comparable to that there was a significant mortality benefit in the ARB group of 46%. observed with other proven secondary prevention measures, such ELITE II trial further evaluated this benefit in more than 3000 as aspirin, beta blockers, and lipid-lowering agents. The institution patients with heart failure and an ejection fraction < 40%. 162 This of therapy should occur early with blood pressure parameters and trial failed to show a benefit of losartan in reducing mortality or close monitoring of hemodynamics. The choice of agent seems morbidity, with a slight nonsignificant benefit for the ACE unimportant, but the dose should be maximized as tolerated by the inhibitor group. patient (see Table 11-1).

ANGIOTENSIN RECEPTOR BLOCKERS

block the end pathway of the RAAS, the AT 1 receptor. All of the In the subset of patients not receiving an ACE inhibitor (7%), potentially damaging cardiovascular actions of A II are mediated through this receptor, including vascular and cardiac hypertrophy, valsartan. A similar reduction of car diovascular mortality or aldosterone secretion, and vasoconstriction. 157 As described in an heart failure was once again noted in CHARM-Alternative with earlier section, A II can also be generated by non-ACE-mediated the use of candesartan 164 (see Table 11-4). pathways, such that inhibition of this enzyme will not lead to complete cessation of A II synthesis. The pharmacologic rationale for interruption of the RAAS with an ARB includes a decrease in AT 1 receptor activation that mediates many of the negative The quinine-related side effects of cough and angioedema are cardiovascular effects and an increase in circulating A II, which common with ACE inhibitors, affecting approximately 5% to undergoes a compensatory rise during ARB therapy that activates 10% of patients, particularly women, African Americans, and the AT 2 receptor, providing additional benefit.

Role of Angiotensin Receptor Blockers

Post-Myocardial Infarction

ACE inhibitors have been shown to reduce the risk of death and cardiovascular events after acute myocardial infarction. Two studies have assessed whether ARBs offer benefits equivalent or superior to ACE inhibitors after myocardial infarction, OPTIMAAL and VALIANT. The Optimal Trial in Myocardial Infarction with the Angiotensin II Antagonist Losartan (OPTIMAAL) evaluated the Renal Effects benefit of losartan compared with captopril in 5477 patients with AT 1 receptor antagonism has been shown to improve acute myocardial infarction complicated by heart failure. 158 No significant difference was noted in all-cause mortality, cardiac mortality, or reinfarction, with losartan unable to achieve noninferiority. There was, however, a trend in event rates in favor of captopril and significantly fewer cardiovascular deaths with captopril (13.3% versus 15.3%). A lower dose of ARB in OPTIMAAL than that suggested in other ARB trials such as LIFE and RENAAL probably contributed to these results. Higher doses of losartan result in an increase in plasma renin activity and A II concentration, indicating a more potent negative feedback at the highest doses. 159 The Heart Failure Endpoint Evaluation of A II Antagonist Losartan (HEAAL) study confirmed the dose effect of losartan, with 150 mg compared with 50 mg showing a significant 10% reduction in Valsartan (MARVAL) in patients with type 2 diabetes mellitus, mortality and morbidity in heart failure patients intolerant to ACE

The Valsartan In Acute Myocardial Infarction (VALIANT) $\frac{17.4}{1}$ (n = 332) with type 2 diabetes and microalbuminuria treated with evaluated the effect of the ARB valsartan versus captopril alone or in combination in 14,703 patients with acute

myocardial infarction complicated by left ventricular systolic 173 dysfunction or heart failure. 160 The population of patients studied was identical to that enrolled in the ACE inhibitor trials after myocardial infarction, SAVE, AIRE, and TRACE, and the dose of valsartan used showed more blood pressure lowering effects than were seen with low-dose losartan in the OPTIMAAL trial.

failure hospitalization was similar between the groups, around 20%. ARBs were noninferior to ACE inhibitor therapy and achieved the same benefit as ACE inhibitors noted in their respective trials.

Congestive Heart Failure

heart failure patients was the Evaluation of Losartan In The Elderly (ELITE) trial. 161 This trial compared the potential renal benefits of losartan with captopril in 722 elderly patients with heart failure. Whereas no benefit in renal function was observed in either arm,

In contrast to ELITE II, the Val-HeFT trial compared the ARB valsartan with standard heart failure therapy. 163 The majority of patients enrolled (93%) in Val-HeFT were already receiving an ACE inhibitor. There was no mortality benefit, but a significant reduction was shown in the combined end point of morbidity Despite the therapeutic success of ACE inhibitors, ARBs emerged to and mortality driven primarily by heart failure hospitalizations. there was a substantial 33% mortality benefit associated with

Kinin-Mediated Side Effects

Asians. 165,166 The use of ARBs in patients who are ACE inhibitor intolerant has been effective with minimal ACE-related adverse effects. ¹⁶⁷ However, blocking of the AT ₁ receptor with selective antagonists is known to increase plasma levels of A II. A II can then interact with other angiotensin receptors including AT 2, stimulating bradykinin release through increased kininogen activation. Cough and angioedema, mediated by bradykinin production, are less common with ARBs but not totally eliminated. 168

albuminuria and to reduce progression of renal disease. The RENAAL and IDNT trials, as described, evaluated the long term benefits of ARB therapy in patients with proteinuria and type 2 diabetes. 132,133 In RENAAL, treatment with losartan for an average of 3.4 years was associated with a 25% reduction of risk for doubling of serum creatinine and a 28% reduction in the risk for development of ESRD, independent of its blood pressurelowering effect. IDNT showed a 20% reduction in the composite endpoint of doubling of creatinine, development of ESRD, or death with irbesartan compared with placebo and 23% reduction compared with amlodipine, for an average of 2.6 years.

Another similar trial, Microalbuminuria Reduction with showed an improvement in proteinuria in patients

valsartan instead of amlodipine. 169 The primary end point of percentage change in urine albumin excretion rate was significantly lower after 24 weeks of treatment for ARB (56%) than for CCB (92%). Valsartan lowered proteinuria in both hypertensive and normotensive subgroups, with more patients reverting to normoalbuminuria with the short-term ARB (30% versus 14.5%).

In the Telmisartan Randomized Assessment Study in ACE Intolerant Subjects with Cardiovascular Disease (TRAN I SCEND)

trial, 5927 patients with known cardiovascular disease or diabetes signaling, and cardiomyocyte apoptosis while improving left without evidence of macroalbuminuria and intolerant to ACE inhibitor ventricular systolic and diastolic dysfunction, chamber dila tion, and were treated with either telmisartan or placebo for a mean of 56 months. pulmonary congestion. 175 Furthermore, aliskiren improves nitric At base -11 line, only 10% of patients had evidence of oxide availability and has minimal effect on platelet activity or microalbuminuria. There was no difference in the composite renal coagulation, except for a moderate increase in antithrombin III. 176,177 outcome of dialysis or doubling of creatinine, changes in GFR, and changes in albuminuria. Albuminuria, however, was less likely to and sustained reduction in blood pressure when it is used as progress with ARB therapy.

blockade of the AT₁ receptor prevents the progression of proteinuria effects of hypertension over time and is a predictive marker of and renal decline. The benefit in high-risk patients without cardiovascular events in patients with CAD risk factors. The significant albuminuria is less well evidenced. ARBs, like ACE Aliskiren in Left Ventricular Hypertrophy (ALLAY) trial assessed inhibitors, are renoprotective, particularly in patients with type 2 whether aliskiren mono therapy or combination therapy with an diabetes and diabetic nephropathy (see Table 11-2).

Summary

ARBs are an effective alternative to ACE inhibitors in patients after myocardial infarction or with heart failure. Increased doses are needed to achieve their maximum benefit for clinical outcomes and improved heart failure symptoms. ARBs may be used for patients who are ACE inhibitor intolerant, but the risk of kinin-mediated side the reductions in left ventricular mass with DRIs were similar to effects is not totally eliminated (see Table 11-1).

DIRECT RENIN INHIBITORS

The RAAS can be inhibited at different levels of the pathway. Despite treatment with ACE inhibitors and ARBs, high rates of cardiovascular events in individuals with CAD and heart failure and high rates of progression to renal failure in individuals with diabetic nephropathy persist. There is growing evidence that these drugs may not be able to optimally inhibit the RAAS because of their inability to block alternative pathways of A II production, the activation of feedback mechanisms that result in increased plasma renin concentration and plasma renin activity, and the stimulation of the prorenin receptor pathway.

In addition, intracellular A II may not be blocked by ACE inhibitors or ARBs as these drugs are unable to enter the intracellular space. Blocking of RAAS directly at the renin level, the rate-limiting step of A II production, neutralizes any compensatory increase in plasma renin activity and prevents the formation of both AI and A II, allowing more complete RAAS suppression. 98 Decreasing plasma renin activity, perhaps the most accurate reflection of the level of RAAS activation, may further reduce the potential harmful effects of the RAAS, providing improved end-organ protection. Cer tainly, there appears to be cardiovascular benefit associated with lower plasma renin activity.

Role of Direct Renin Inhibitors

Aliskiren is a potent nonpeptide renin inhibitor that because of its high specificity has a low drug interaction profile. ¹⁷¹ It functions by binding to the enzymatic site of renin and rendering it inactive. Aliskiren binds to plasma renin before the uptake by the tissue prorenin receptor, but it is unclear whether renin inhibitors could inhibit receptor-bound, non-proteolytically activated prorenin or modify the structure of renin, altering its ability to interact with the prorenin receptor. 172 No observed changes have been shown in either renin- or prorenin-mediated prorenin receptor activation with direct renin inhibitors (DRIs), indicating that aliskiren does not affect its binding site for the prorenin receptor. ¹⁷³

The effects of DRIs are well established in animal studies. Aliskiren has been shown to prevent atherosclerosis progression compared with other antihypertensive agents with similar blood pressure-lowering effect. ¹⁷⁴ It reduces fibrous cap thinning, creates less vulnerable plaque with decreased inflammation and lipid content, and prevents media degeneration. In untreated animals, evidence of advanced plaque with a large necrotic lipid core was found in more than 80% of cases compared with only 25% for treatment with aliskiren. Renin inhibition has also been shown to attenuate infarction-related changes, the indices of A II intracellular

In clinical trials, aliskiren provides an effective dose dependent monotherapy or in combination with other anti hypertensive agents. In the treatment of the hypertensive and diabetic patient, 178-181 Left ventricular hypertrophy represents a manifestation of the ARB reduced left ventricular mass in overweight, hypertensive patients with left ventricular hypertrophy. 182 The combination of a DRI with ARB did not show a statistically significant difference in the reduction of blood pressure or left ventricular mass during a 9month follow-up period. Whether the lack of combination effect can be attributed to a lower baseline blood pressure or left ventricular mass compared with other trials, short duration of therapy, or altered renin states among these patients remains unclear. However, those achieved with an ARB with equivalent tolerability.

Other surrogate marker trials looking at the effects of DRI in heart failure and albuminuria as well as in myocardial infarction and left ventricular remodeling have been performed. The Aliskiren Observations of Heart Failure Treatment (ALOFT) trial showed that aliskiren, added to optimal medical therapy with an ACE inhibitor or ARB and beta blocker in all patients and an aldosterone receptor antagonist in approximately one third, was associated with improvements in the heart failure markers brain natriuretic peptide (BNP), N-terminal pro-BNP (NT-proBNP), and urinary aldosterone levels. 183 There was a fivefold greater decrease in BNP levels compared to placebo. Reductions in BNP and NT-proBNP have been noted with other RAAS inhibitors and are associated with improved cardiovascular outcomes.

The Aliskiren Study in Post-MI Patients to Reduce Remodeling (ASPIRE) trial assessed the effects of DRI on left ven tricular remodeling in 820 patients with ejection fractions less than 45% compared with placebo when added to standard therapy including an ACE inhibitor or ARB. 184 After 36 weeks of therapy, the addition of aliskiren within the 7 to 42 days after a myocardial infarction showed no significant improvement in left ventricular end-systolic volume or com posite endpoint of cardiovascular death, CHF hospitalization,

however, a higher rate of hyperkalemia, hypotension, and renal decrease in CHF hospitalizations by 35% with the use of dysfunction. Additional trials like Aliskiren and Valsartan to Reduce spironolactone after 2 years. From RALES, the benefit of NT-proBNP via Renin-Angiotensin-Aldosterone System Blockade mineralocorticoid receptor blockade in subjects with CHF can be (AVANTE GARDE) will further evaluate the potential role of noted beyond RAAS inhibition and the diuretic effects of aliskiren after an acute myocardial infarction.

Diabetes Using Cardiovascular and Renal Disease Endpoints which is now considered a standard of care. (ALTITUDE) and Aliskiren Trial to Mediate Outcomes Prevention in Heart Failure (ATMOSPHERE). 185 The ALTITUDE trial is Failure Efficacy and Survival Study (EPHESUS), the benefit of designed to evaluate the benefit of adding aliskiren to conventional adding eplerenone to optimal medical therapy for the treatment of medical therapy that includes either an ACE inhibitor or ARB long heart failure after acute myocardial infarction was evaluated. 190 term in high-risk diabetic patients. It is an event-driven study EPHESUS enrolled 6600 subjects with an ejection fraction of 40% or looking at both cardiovascular and renal outcomes. The less, and treated with coronary artery fusion therapy if needed, 3 to ATMOSPHERE trial will evaluate whether aliskiren can delay the 14 days after an acute myocardial infarction. Subjects were time to cardiovascular death or CHF hospitalization in patients with randomized to either eplerenone or placebo; results showed a heart failure (see Table 11-1).

Summary

Aliskiren is an effective and well-tolerated DRI for patients with hypertension. It can be added to multiple therapies with proper monitoring of electrolytes and renal function. Surro gate marker trials have suggested significant potential in the management of patients with heart failure and nephropathy. The results of future large-scale outcome trials will help elucidate the clinically relevant protective end-organ effects of this class of medication. For now, in patients who are able to tolerate the older RAAS blockers, ACE inhibitors, or ARBs, there is no reason to prefer aliskiren. Aliskiren can be used in patients who cannot tolerate either an ACE inhibitor or an ARB. Although morbidity and mortality data are lacking, there is good evidence that combination therapy can improve blood pressure control, and ongoing randomized trials are looking at potential further event reduction with this type of combination therapy (see Table 11-1).

ALDOSTERONE RECEPTOR ANTAGONISTS/MINERALOCORTICOID RECEPTOR

Early results from the CONSENSUS trial showed that in addition to sympathetic activation and increases in A II, plasma aldosterone levels predicted subsequent cardiovascular morbidity and mortality. 186 This suggested a possible role of aldosterone in the progression of systolic dysfunction. ACE inhibitor or ARB therapy Summary has been shown to incompletely suppress aldosterone production, and an increase can be noted in up to 40% of patients with CHF. 187 Aldosterone is controlled not only by A II but also by potassium levels. 84 With a fall in A II, an increase in potassium will stimulate aldosterone production.

Role of Mineralocorticoid Receptor Blockers

The two agents used for aldosterone receptor blockade are spironolactone, the older of the two, and eplerenone. Eplerenone is 100 times more selective for the aldosterone receptor and has limited further stimulate aldosterone production . No evidence is affinity for androgen and progesterone receptors. 188 There is a lower currently available that shows a benefit of aldosterone blockade incidence of gynecomastia and breast pain in men and menstrual in CAD patients without CHF (see Table 11-4). irregularities in women with eplerenone compared with spironolactone (< 1% versus 10%). The incidence of hyperkalemia is approximately 3% to 6% for spironolactone and 1% to 3% for eplerenone. The short half-life of eplerenone and lack of active metabolites may lessen the risk of hyperkalemia. 189

The Randomized Aldactone Evaluation Study (RALES) 175 further evaluated the benefit of adding an aldosterone receptor antagonist to optimal medical therapy. 86 RALES enrolled 1663 subjects with NYHA Class III or Class IV heart failure with an ejection fraction of 35% or less while receiving treatment with an ÁCE inhibitor and a diuretic. Patients received either spironolactone

or ejection fraction reduction of greater than 6%. There was, or placebo, and results showed a decrease in mortality by 30% and a aldosterone blockade. However , one of the limitations of this Ongoing outcomes trials include Aliskiren Trial in Type 2 important study was the virtual absence of beta blocker therapy,

> In the Eplerenone Post-Acute Myocardial Infarction Heart reduced all-cause mortality by 15% and cardiovascular death or hospitalization for cardiovascular events by 13%.

The difference in risk reduction compared with RALES can potentially be attributed to subjects with a slightly improved baseline ejection fraction, successful reperfusion therapy after myocardial infarction, recovery of ventricular stunning, and improved use of beta blockade, which was 75% compared with only 12% in RALES. Nevertheless, this study expanded the use of aldosterone blockade to include post-myocardial infarction patients with impaired systolic function. Aldosterone blockade should be initiated in this population between days 3 and 14, when patients are hemodynamically stable after starting an ACE inhibitor or ARB and beta blocker therapy.

A meta-analysis evaluating 19 trials of aldosterone antagonists in CHF and post-myocardial infarction showed that aldosterone blockade reduced all-cause mortality by 20% (25% in 14 CHF trials and 15% in 4 myocardial infarction trials). 191 The trials with longer duration of follow-up were associated with the greatest reduction in all-cause mortality. From RALES and EPHESUS, aldosterone blockade has a clear mortality benefit with and without acute myocardial infarction in heart failure above standard therapy with an ACE inhibitor. Proposed mechanisms include improved endothelial function with a subsequent reduction in acute coronary events, decrease in myocardial fibrosis with a reduction in arrhythmogenic events, maintenance of potassium and magnesium levels, and improvements in fibrinolytic balance due to the direct effect of aldosterone on PAI-1 192 (see Table 11-1).

Mineralocorticoid receptor blockers (MRBs) have an important role in the treatment of heart failure, especially after an acute myocardial infarction, above standard therapy with ACE inhibition. It would seem logical that these same mechanisms and effects of aldosterone receptor antagonism, when applied to patients with CAD but no heart failure, would be evident. In addition, CAD patients without CHF would be more likely to tolerate aldosterone blockade, given their decreased likelihood of renal insufficiency and increased use of diuretics, which may

176 RAAS COMBINATION THERAPY

Despite the many therapies described, a significant number of patients with cardiac and renal disease continue to progress. Incomplete blockade of the RAAS is a potential explanation for these results. Given the multiple feedback loops and alternative pathways described within the RAAS, inhibition of the RAAS at one level leads to compensatory activation at another level, which can diminish the intended therapeutic effect (see Fig. 11-2 and Table 11-3). Strategies to improve the efficacy of RAAS blockade include the combination of ACE inhibitors and ARBs. In addition, that may offer novel 11 ways to suppress the RAAS.

potential means for enhanced suppression. 193 ACE inhibition noted with a reduction in systolic blood pressure of 2 to 3 mm Hg. reduces the levels of A II and increases A I. Increased AI may stimulating non-AT 1 receptors with unclear effects.

regardless of the sodium intake. Because the augmentation of failure despite ACE inhibitor therapy (see Table 11-4). bradykinin levels may also contribute to the net therapeutic

ways of angiotensinogen-A I may limit their effectiveness.

Incomplete RAAS blockade or compensatory RAAS activation limits the potential efficacy of any one of these protection compared with either medication alone.

Dual RAAS Blockade

ACE Inhibitor/ARB

The Val-HeFT and the Candesartan in Heart Failure (CHARM- Added) trials evaluated 5010 and 2548 patients, respectively, with NYHA Class II-IV heart failure and a left ventricular ejection fraction of less than 40% who were being treated with an ACE inhibitor. 163,197 Patients were assigned to an ARB, either valsartan or candesartan, or placebo and were observed between 23 and 41 months. In Val-HeFT, overall mortality was similar in the two groups, with a 24% reduction in heart failure hospitalization. In CHARM-Added, there was a 15% significant reduction in cardiovascular mortality or heart failure admissions. The apparent differences between the Val-HeFT and CHARM-Added trials can be potentially explained by the choice of ARB. Nevertheless, in patients with symptomatic heart failure, there is some added clinical benefit in terms of reducing the morbidity of cardiovascular disease by adding an ARB to ACE inhibitors, with a potential improvement in cardiovascular mortality.

In the VALIANT trial, the combination of valsartan and captopril showed no incremental benefit of ARB added to an ACE inhibitor in acute myocardial infarction complicated by heart failure. 160 This lack of benefit was noted in all-cause mortality and in the secondary endpoint of cardiovascular death, recurrent myocardial infarction, or heart failure hospitalization, despite additional lowering of blood pressure of 2.2 mm Hg. In the Ongoing Telmisartan Alone and in Combination with Ramipril Global Endpoint Trial (ONTARGET), 25,620 patients who had vascular disease or high-risk diabetes without heart failure, with nearly three quarters having CAD, were randomized to ramipril, telmisartan, or a combination of both. 198 No

increased understanding of the role of aldosterone in vascular significant difference in overall mortality or rates of secondary progression and the availability of DRIs allow varied regimens outcomes including a composite of cardiovascular death, myocardial infarction, or stroke was noted except for a slight Understanding of the differential effects of each therapy on increased risk of renal impairment in the combination ACE the components of the RAAS is important to understand a inhibitor/ARB group. This lack of benefit, as in VALIANT, was also

The results of VALIANT and ONTARGET are discordant with be converted to A II through non-ACE-dependent pathways, those of trials like Val-HeFT and CHARM-Added. The differences also known as the ACE escape. ARBs block the angiotensin I in cardiovascular risk between patients with stable heart failure and AT 1 receptor, causing an increase in both AI and A II. Elevated patients with acute myocardial infarction, higher risk of early death levels of A II may potentially compete with the ARB for the AT and myocardial infarction, may account for the differences seen $_1$ receptor, maintaining AT $_1$ receptor signaling , and between these trials. In the latter trials, the ARB is started on a background of ACE inhibitor, not concurrently, and their doses are Both ACE inhibitors and ARBs suppress the negative feed not titrated to proven efficacy levels. Less than half of patients were back inhibition of renin release and cause a compensatory rise receiving full-dose ACE inhibitor therapy, and decisions about the in the plasma renin concentration and plasma renin activity. dose and choice of an ACE inhibitor were left to individual 194 In addition, there is a notable secondary rise in aldosterone, physicians. Furthermore, all of these patients had symptomatic heart

Renal Effects. Whereas ACE inhibitors and ARBs have been benefits of ACE inhibition, the combination of an ARB that shown to be renoprotective when used alone, the effect of their stimulates bradykinin production with an ACE inhibitor that combined use on the progression of renal disease is less certain. inhibits bradykinin degradation may provide a theoretical Secondary endpoints from the ONTARGET trial showed that there basis for augmentation of the bradykinin-nitric oxide was an increase in the composite endpoint of dialysis, doubling of serum creatinine, or death of 14.5% with ramipril/telmisartan MRBs increase all components of RAAS: plasma renin compared with ACE inhibitor or ARB alone (13.5%). 199 This activity, plasma renin concentration, and AI, A II, and difference in renal endpoint was driven primarily by dialysis aldosterone levels. 195,196 DRIs, on the other hand, lead to a treatment for episodes of acute renal failure, not by the development decrease in plasma renin activity despite a compensatory of ESRD, and there was no significant difference in the incidence of $increase\ in\ plasma\ renin\ concentration.\ ^{194}\ Renin\ inhibitors\ can\ creatinine\ doubling\ .\ Subgroup\ analysis\ showed\ that\ dual\ blockade$ antagonize the increase in plasma renin concentration induced was harmful only in individuals with low renal risk, without by the ACE inhibitor, ARB, or MRB; however, nonrenin path diabetes and albuminuria. This increased event rate was matched by a 7% reduction in urinary albumin excretion compared with ACE inhibitor therapy.

The Candesartan and Lisinopril Microalbuminuria (CALM) trial classes of medications on cardioprotection and renoprotection. evaluated 199 patients with microalbuminuria, hypertension, and Combinations of these medications or the addition of other - type 2 diabetes with 12 weeks of candesartan, lisinopril, or antihypertensive medications, specifically CCBs, may provide combination therapy. 200 There was a significant reduction in urinary more complete RAAS suppression and better end-organ ACR with ACE inhibitor/ARB compared with ACE inhibitor or ARB alone (50% versus 39% versus 24%). However, the mean reduction in blood pressure (systolic blood pressure, 25 mm Hg versus 17 mm Hg and 14 mm Hg) was significantly greater with dual versus mono therapies. Dual ACE inhibitor/ARB blockade in CALM was shown to reduce proteinuria, but it may have been explained by improved blood pressure control.

> In the Valsartan in Combination with Lisinopril in Hypertensive Patients with Microalbuminuria (VALERIA) trial, 133 patients with hypertension and microalbuminuria were evaluated by ACE inhibitor/ARB combination therapy compared against high-dose ACE inhibitor or ARB monotherapy. 201 After 30 weeks of treatment, there was a 34% reduction in

with no significant difference compared with the ARB group. 11-3). Hypotension was the most frequent adverse event, occurring at 9.3% for ARB and 11.6% for ACE inhibitor/ARB.

combination therapy (see Table 11-2).

therapy can be noted in patients with uncontrolled, symptomatic outcomes and in turn potentially cardiac outcomes (see Table 11-2). heart failure. However, there is no role for ACE inhibitor/ARB combination therapy in patients with stable CAD or in secondary prevention without heart failure. The utility of combination therapy must be weighed against the increased risk of hypotension, hyperkalemia, and renal function decline (see Table 11-1).

ACE Inhibitor/MRB or ARB/MRB

inhibitor or MRB/ARB in patients with left ventricular dysfunction improved ATP-sparing abilities and t-PA/PAI-1 balance can after myocardial infarction or heart failure. Evidence of this benefit potentially work synergistically for enhanced cardioprotection. The was noted in their respective trials, EPHESUS and RALES, and in a vasodilation caused by CCBs also stimulates both the RAAS and the meta-analysis predominantly powered by these trials. An overall all-sympathetic nervous system, which can lead to reflex cause mortality benefit of 20% with MRB/ACE inhibitor or vasoconstriction and tachycardia. ACE inhibition can counteract MRB/ARB (25% improvement in patients with CHF and 15% this effect. 36 improvement in patients with acute myocardial infarction) can be

inflammatory state, ²⁰² alterations in fibrinolysis, ²⁰³ stimulation of randomized to either ACE inhibitor/ CCB therapy of A II receptors. ²⁰⁶ By blocking aldosterone at its receptor, MRBs (atenolol/bendroflumethiazide). The trial was stopped early may attenuate the effects of aldosterone on the kidney, reducing because of an ~35% reduction in CHD events at a mean follow-up of proteinuria and renal outcomes. Several small studies have about 3.3 years; there was also an 11% difference in all-cause addressed the additive benefit of MRB with an ACE inhibitor or mortality noted between the two groups at 5.5 years. Secondary ARB. Epstein and colleagues ²⁰² evaluated the effect of low doses (50 endpoints included a 24% decrease in cardiovascular mortality, a company events and a 23% difference in stroke. to 100 mg/day) of eplerenone added to enalapril on urinary ACR in 13% decrease in all coronary events, and a 23% difference in stroke. patients with type 2 diabetes. A nearly 50% reduction in albumin significant hyperkalemia. 207

Mehdi and coworkers ²⁰⁸ assigned 81 patients with hypertension rather than ARB to maximal AČE inhibitor therapy affords greater inhibitor/CCB therapy, with a 46% reduction in all-cause mortality. renal protection in patients with diabetic nephropathy. 178 The International Verapamil-Trandolapril Study Trial (INVEST) Nevertheless, the long-term effect of ACE inhibitor/MRB on renal outcomes has yet to be determined. In patients with persistent proteinuria despite maximal doses of ACE inhibitor or ARB, lowdose MRB could be added to reduce proteinuria with monitoring for hyperkalemia (see Table 11-2).

ACE Inhibitor/DRI or ARB/DRI

Evidence for dual blockade with a renin inhibitor and ACE inhibitor or ARB is still minimal. In the ALOFT trial, the DRI aliskiren, when added to an ACE inhibitor or ARB and an aldosterone receptor antagonist in approximately one third, was associated with a significant improvement in intermediate endpoints of heart failure and BNP and urinary aldosterone levels. This was noted despite an

urine ACR with combination therapy compared with mono therapy. absence of effect on blood pressure. Clinical outcomes for 177 In the ACE inhibitor/ARB group, 38% had normalization of combination DRI/ACE inhibitor or DRI/ARB in patients with microalbuminuria compared with 17% in the ACE inhibitor group, cardiovascular disease remain to be evaluated (see Tables 11-1 and

Renal Effects. In the Aliskiren in the Evaluation of Proteinuria in Diabetes (AVOID) trial, the renoprotective effects of combining Combined therapy with ACE inhibitor and ARB results in more aliskiren with the high-dose ARB losartan, which was shown to have complete blockade of RAAS and reduction or normalization of renoprotective effects in the RENAAL study, was evaluated in microalbuminuria than with high-dose mono therapy. ACE patients with hypertension, diabetes, and nephropathy. 209 Aliskiren, inhibitor/ARB therapy in both diabetic and nondiabetic when added to an ARB and optimal antihypertensive treatment, nephropathies significantly reduces proteinuria. This benefit occurs provided a significant 20% further reduction in urinary ACR. Nearly with a slightly higher incidence of adverse events. Patients at low twice as many patients in the aliskiren group also achieved a renal risk, nondiabetics, and patients without evidence of reduction in urinary ACR of at least 50% from baseline. These proteinuria are at higher risk for worsened renal outcomes with renoprotective effects were independent of the effect of DRI on blood pressure. Reductions in albuminuria in patients with diabetes Summary. The benefit of combination ACE inhibitor/ARB and nephropathy are associated with improvements in renal 11

ACE Inhibitor/CCB Combination Therapy

CCBs counteract excess calcium entry through the voltage- and receptor-operated calcium channels of vascular smooth muscle. ACE inhibitors reduce the vasoconstrictive properties of A II, indirectly through sodium and water balance and directly by reducing interpol calcium release. Smooth muscle was diletted by reducing internal calcium release. Smooth muscle vasodilation by Favorable outcomes were noted by dual blockade with MRB/ACE calcium maintenance and other anti-ischemic properties including

The rationale for fixed-combination ACE inhibitor and CCB & noted in the appropriate patients, NYHA Class III-IV heart failure therapy comes directly from the results of ASCOT, the first trial to with an ejection fraction < 35% or acute myocardial infarction with an ejection fraction < 40% (see Table 11-4).

morbidity between two antihypertensive regimens. 67 ASCOT morbidity between two antihypertensive regimens. 10 ASCOT morbidity between two antihypertensive regimens. 11 ASCOT morbidity between two antihypertensive regimens. 12 ASCOT morbidity between two antihypertensive regimens. 13 ASCOT morbidity between two antihypertensive regimens. 14 ASCOT morbidity between two antihypertensive regimens. 15 ASCOT morbidity between two antihypertensive regimens. 16 ASCOT morbidity between two antihypertensive regimens. 16 ASCOT morbidity between two antihypertensive regimens. 17 ASCOT morbidity between two antihypertensive regimens. 18 ASCO ejection fraction < 40% (see Table 11-4). morbidity between two antihypertensive regimens. ⁶⁷ ASCOT **Renal Effects.** Aldosterone may promote renal fibrosis and renal included 19,257 hypertensive patients with at least three dysfunction by multiple mechanisms, including an increased cardiovascular risk factors but no cardiac disease. Patients were a inflammatory state, 202 alterations in fibrinolysis, 203 stimulation of randomized to either ACE inhibitor/ CCB therapy addressed the additive benefit of MRB with an ACE inhibitor or mortality noted between the two groups at 5.5 years. Secondary

In ASCOT, stable angina was an exclusion criterion for this 5 excretion was noted from baseline as early as 4 weeks and continued study, making assessment of combination therapy on stable CAD for up to 12 weeks. Low-dose MRB demonstrated a similar reduction difficult. However, there was a noted decrease in coronary events in proteinuria compared with high-dose MRB (200 mg/day) without and a 32% reduction in unstable angina. The reduction of coronary and stroke events cannot be entirely explained by a reduction in 9 blood pressure. ²¹⁰ Another trial, EUROPA, conducted in stable CAD , diabetes, and macroalbuminuria who were taking maximal doses patients, showed that the ACE inhibitor perindopril significantly of ACE inhibitor to either ARB (losartan, 100 mg/day) or reduced the composite endpoint of cardiovascular death, spironolactone (25 mg/day). After 48 weeks of therapy, the urine myocardial infarction, and cardiac arrest. Post hoc analysis of this ACR was significantly lower in patients taking ACE inhibitor/MRB trial evaluated the effect of ACE inhibition in 2122 patients receiving (34%) versus ACE inhibitor/ARB (17%). Blood pressure did not CCB, 18% of the patients enrolled. The event rate for major cardiac differ between the groups, suggesting that the addition of MRB events and mortality was reduced by 35.5% with combination ACE

> compared ACE inhibitor/CCB therapy (verapamil/ trandolapril) with beta blocker/diuretic therapy (atenolol/ hydrochlorothiazide) in 22,576 hypertensive patients with CAD. 211 The trial showed that ACE inhibitor/CCB was a welltolerated strategy for treatment. It failed, however, to show a benefit of ACE inhibitor/CCB over beta blocker/diuretic combination therapy. In light of ASCOT, this may suggest that the effect of CCB in combination with an ACE inhibitor may not be due to a pure class effect.

In the Avoiding Cardiovascular Events through Combination Therapy in Patients Living with Systolic Hypertension (ACCOMPLISH) trial, 11,506 hypertensive patients

at high risk for cardiovascular events were randomized to either ACE 11 inhibitor/CCB (benazepril/amlodipine) or ACE inhibitor diuretic (benazepril/hydrochlorothiazide). There was a 20% risk reduction in the composite endpoint of cardiovascular death, nonfatal myocardial infarction or stroke, hospitalization for revascularization. This trial showed that the combination of ACE (Kostis JB, Wilson AC, Freudenberger RS, et al: Long-term effect of diuretic-based therapy on fatal outcomes in enhanced the combination of ACE). inhibitor/CCB is superior to ACE inhibitor/diuretic in reducing cardiovascular events and death in high-risk patients. The 15. Major outcomes in high-risk hypertensive patients randomized to angiotensin converting enzyme difference in systolic blood pressure was less than 1 mm Hg between the two groups, suggesting that the difference in benefit 16. was not related to blood pressure lowering effects. Overall, the ACE inhibitor/CCB combination is an effective therapeutic option 17. Brown MJ, Palmer CR, Castaigne A, et al: Morbidity and mortality in patients randomized to to optimize the management of high-risk patients with hypertension and patients with CAD (see Table 11-1).

Renal Effects. Individually, ACE inhibitors show uniform 18. efficacy in reducing proteinuria and limiting the progression of renal disease. CCBs, as described, have more varied results. Whether this is related to the class of CCB, the underlying renal disease, or the duration of treatment has vet to be determined. Limited information is available for the effect of 20. ACE inhibitor/CCB therapy on proteinuria and renal function, but it appears to confer an advantage. ²¹² Fogari and associates ²¹³ compared the long-term effect of amlodipine and 22. Loscalzo J: Antiplatelet and antithrombotic effects of organic nitrates. Am J Cardiol 70:18B, 1992. fosinopril on urine albumin excretion in 453 hypertensive, 23. diabetic patients. In a 4-year follow-up, ACE inhibitor/CCB provided a 67% reduction in urine albumin excretion compared with monotherapy with ACE inhibitor (46%) or renal impairment reduction in amlodipine/perindopril group versus beta blocker/diuretic. 26. 67 These studies strengthen the recommendation for ACE inhibitor/CCB combination therapy for renoprotection in 27. GISSI-3: Effects of lisinopril and transdermal glyceryl trinitrate singly and together on 6-week high-risk patients (see Table 11-2).

CONCLUSION

In summary, each class of antihypertensive medications has ²⁹. an established role in the management of hypertensive and high-risk cardiovascular patients. Understanding of the complexity of the RAAS has improved our understanding of the close interplay between cardiovascular and renovascular 31. Heitzer T, Just H, Brockhoff C, et al: Long-term nitroglycerin treatment is associated with disease and ways to improve both cardiovascular and renal outcomes. RAAS inhibitors are the cornerstone for secondary prevention. Newer RAAS inhibitors along with combinations 32. Hirai N, Kawano H, Yasue H, et al. Attenuation of nitrate tolerance and oxidative stress by an of different RAAS inhibitors and RAAS inhibitors with CCBs may provide more complete RAAS suppression and improve cardioprotection and renoprotection.

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Cholesterol/Dyslipidemia

CHAPTER 12

Evaluation and Management of Dyslipidemia in Children and Adolescents

KEY POINTS
Christian D. Nagy and Peter O. Kwiterovich, Jr.

- Early lesions of atherosclerosis begin in childhood and are related to antecedent CVD risk factors. Environmental factors such as diet, obesity, and exercise and certain inherited dyslipidemias influence the progression of such lesions. Detection of youth at risk for atherosclerosis includes an integrated evaluation of these predisposing factors.
- Treatment starts with weight control, exercise, and a diet low in saturated fat, *trans*-fat, and cholesterol, supplemented with water-soluble fiber, plant stanols, or plant sterols.
- Drug therapy, especially with HMG-CoA reductase inhibitors or bile acid sequestrants, can be considered in those with a positive family history of premature CVD and LDL-C >160 mg/dL, after a period of dietary and hygienic measures.

 Candidates for drug therapy often include those with familial hypercholesterolemia, familial combined hyperlipidemia, the metabolic syndrome, polycystic ovarian syndrome, type 1 or 2 diabetes, and the nephrotic syndrome.
- Dietary and drug therapy appears to be safe and efficacious.

 Early identification and treatment of children with CVD risk factors and dyslipidemia are likely to retard the atherosclerotic process.

Disorders of lipid and lipoprotein metabolism are characterized by abnormalities of the major lipoprotein classes, chylo microns, verylow-density lipoproteins (VLDL), low-density lipoproteins (LDL), and high-density lipoproteins (HDL). Dys lipidemia can result from the expression of a mutation in a single gene that plays a significant role in lipoprotein metabolism, reflect the effects of multiple genes, or be caused by environmental influences such as excessive dietary intake of fat and calories and limited physical activity, particularly in association with overweight or obesity.

The major clinical complication of dyslipidemia is a predisposition to atherosclerosis starting early in life and leading to cardiovascular disease (CVD) in adulthood. Multiple trials in adults have demonstrated that reduction in the plasma levels of LDL results in the decreased prevalence of coronary artery disease (CAD) and stroke. Low ering of LDL also produces a decrease in the angiographic progression of CAD and even modest regression in some In children with familial hypercholesterolemia, a decrease in LDL reduces carotid intima media thickness (CIMT) and lipid screening recommendations have recently been

 Optimal identification and treatment of high-risk youth requires an integrated universal screening, evaluation, treatment, and follow-up program.

updated by the American Academy of Pediatrics. ¹

This chapter presents a theoretical and practical approach to the diagnosis and management of dyslipidemia in children and adolescents, in the hope of preventing or retarding the progression of atherosclerosis - early in life.

BACKGROUND

A number of studies strongly support the notion that atherosclerosis and CVD risk factors originate in childhood and adolescence and that treatment should begin early in life. ² Several longitudinal pathological studies found that early atherosclerotic lesions of fatty streaks and fibrous plaques in children, adolescents, and young adults who died of accidental deaths are significantly related to higher antecedent levels of total cholesterol (TC) and LDL-cholesterol (LDL-C), lower levels of HDL-cholesterol (HDL-C), and other CVD risk factors, such as higher blood pressure, cigarette smoking, and obesity. ²⁻⁴

Major prospective population studies, including the Muscatine Study, 5.6 the

12

TABLE 12-1 Classification and Properties of Major Plasma Lipoproteins

TABLE 12-1 Classifica	ition and Properties of Major Plasma Lip	oproteins		
	Chylomicrons	Very-Low-Density Lipoproteins	Low-Density Lipoproteins	High-Density Lipoproteins
Hydrated density (g/mL)	< 0.95	0.95-1.006	1,019-1,063	1.063-1.21
Electrophoretic migration	original	Pre -S	AND	A
Molecular weight	50-1000 x 10 a	10-80 X 10 a	2.3 X 10 s	1.8-3.6 x 10 s
Average composition (%)				
Cholesterol*	3	22	50	20
Triglycerides	90	55	5	5
Phospholipid	6	15	25	25
Protein	1	8	20	50
Major apolipoproteins			Apo B-100	
	B-48 Apo AI, IV	Apo B-100 Apo CI, II, III Apo E		Apo AI, II Apo CI, II, III Apo E
original	Bowel	Liver	Product of VLDL catabolism	Liver, intestines
Function	Intestinal transport of triglycerides and cholesterol	Hepatic transport of triglycerides and cholesterol	Provides cholesterol to cells	Reverse cholesterol transport

^{*}Includes the mass of cholesteryl ester and esterified cholesterol.

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Bogalusa Heart Study, ^{7,8} the Coronary Artery Risk Develop ment in Young Adults (CARDIA) study, ⁹ the Special Turku Coronary Risk Factor Intervention Project (STRIP), ¹⁰ and the Cardiovascular Risk in Young Finns Study, ^{11,12} demonstrated that CVD risk factors in children and adolescents, particularly LDL-C and obesity, predicted subclinical manifestations of atherosclerosis in young adults, as estimated by coronary artery calcium, CIMT, or brachial flow-mediated dilation. With respect to clinical manifestations of atherosclerosis, medical students at Johns Hopkins who had a TC level > 207 mg/dL had five times the risk for development of CVD 40 years later than did those students who had a TC level < 172 mg/dL. ¹³

Studies in high-risk youth, selected by virtue of CVD in one parent, demonstrated the familial aggregation of dyslip idemia in such children. For example, half of the young progeny of men with premature CVD before 50 years of age had one of seven dyslipidemic profiles: elevated LDL-C alone (type IIa) or combined with high triglycerides (type IIb); ele vated HDL-C triglycerides alone (type IV); low (hypoalphalipoproteinemia); and type IIa, type IIb, or type IV also accompanied by low HDL-C. 14 Elevated levels of apolipoprotein B, in the presence of normal LDL-C (hyperapobetalipoproteinemia), were prevalent in young with adults CVD offspring of premature hyperapobetalipoproteinemia. 15 The levels of apolipoprotein B (apo B) and apolipoprotein AI (apo AI), the major apolipoproteins of LDL and HDL, respectively, and the ratio of apo B to apo AI in young off spring from Bogalusa were stronger predictors of premature CAD in their parents than the levels of either LDL-C or HDL-C. 16

Inherited lipoprotein disorders that often present in youth at high risk for future CVD include familial hypercholesterolemia (FH), caused by a defect in the LDL receptor (LDLR), and familial combined hyperlipidemia (FCHL) and its metabolic cousin, hyperapobetalipoproteinemia, both prototypes for overproduction of VLDL in the liver. Increased synthesis and secretion of VLDL are often driven by an increased flux of free fatty acids (FFA) from the adipose tissue to the liver, a metabolic abnormality usually accompanied by insulin resistance and the dyslipidemic triad of elevated triglycerides, increased small, dense LDL particles (sLDL-P), and low HDL-C.

In addition, secondary accelerated atherosclerosis requiring aggressive lipid management can be encountered in certain clinical settings early in life, such as in children and adolescents with diabetes mellitus, end-stage renal disease on hemodialysis, or nephrotic syndrome; after renal or heart transplantation; in HIV-positive patients receiving protease inhibitors; in childhood cancer survivors; or in children and adolescents receiving chronic antipsychotic treatment.

LIPID AND LIPOPROTEIN CLASSIFICATION

Cholesterol, triglycerides (TG), phospholipids (PL), and cholesteryl esters (CE) are major lipid classes that are essential components found in dietary fat, plasma lipoproteins, and cells. Lipids are hydrophobic and do not circulate freely in plasma but rather as part of lipid-protein macromolecular complexes called lipoproteins. Plasma lipoproteins are generally spherical particles consisting of a core that contains non polar lipids, mostly TG and CE, surrounded by a surface coating consisting of proteins (apolipoproteins), and polar lipids, such as PL and unesterified (free) cholesterol (FC). Lipoproteins function as transport vehicles for water insoluble lipids and carry them to their sites of metabolism or deposition.

Plasma lipoproteins have been classified by their density and electrophoretic mobility into chylomicrons, VLDL (density < 1.006 g/mL) or pre-beta (pre- Ш) lipoproteins, LDL (density 1.019-1.063 g/mL) or beta (Ш) lipoproteins, and HDL (density 1.063-1.21 g/mL) or alpha (a) lipoproteins (Table 12-1). Intermediate-density lipoproteins (IDL; density 1.006 1.019 g/mL; also called VLDL remnants) are produced after the hydrolysis of the TG on VLDL. Electrophoresis allows the separation of the plasma lipoproteins on the basis of differences in size and electrophoretic charge. After electrophoresis , chylomicrons remain at the origin, and VLDL, LDL, and HDL migrate in the same positions as pre- Ş -, Ş -, and a -globulins, respectively. IDL migrate to a position between the pre- Ş - and Ş -globulins. The hydrated density of the

TABLE 12-2		es of Major Plasma Apolipoprotei	
Apolipoprotein	Molecular Weight	Chromosomal Location	Function
Apo A-I	29,016	11q23	Cofactor LCAT; facilitates both the transfer of cell cholesterol by ABCA-1 to nascent HDL and the delivery of CE and FC on HDL to liver through SR-B1
Apo A-II	17,414	1q21-23	Inhibits TG hydrolysis by HL and VLDL
Apo A-IV	44,465	11q23	Activates LCAT, promotes formation of chylomicrons
Apo A-V	39,000	11q23	Stimulates proteoglycan-bound LPL
Apo B-100	512,723	2p24-p23	Secretion of VLDL from liver; binding ligand to LDLR
Apo B-48	240,800	2p24-p23	Secretion of chylomicrons from intestine
Apo C-I	6630	19q13.2	Inhibits apo E binding to LDLR; stimulates LCAT; inhibits CETP and SR-B1
Apo C-II	8900	19q13.2	Cofactor LPL
Apo C-III	8800	11q23	
			Noncompetitive inhibitor of LPL; inhibits binding of apo E on TG-rich lipoproteins to LDLR
Apo D	19,000	3q26.2	Promotes reverse cholesterol transport
Apo E	34,145	19q13.2	Binding ligand for LRP on chylomicron remnants and VLDL and IDL for LDLR

CE, cholesteryl esters; CETP, cholesteryl ester transfer protein; FC, free cholesterol; HDL, high-density lipoprotein; HL, hepatic lipase; LCAT, lecithin-cholesterol acyltransferase; IDL, intermediate-density lipoprotein; LDL, low-density lipoprotein; LDLR, LDL receptor; LPL, lipoprotein lipase; LRP, LDL-like receptor protein; SR-B1, scavenger receptor B1; TG, triglycerides; VLDL, very-low-density lipoprotein.

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lipoproteins is related to their chemical composition and the relative content of lipid and apolipoprotein.

Chylomicrons contain 99% lipid, mostly TG (see Table 12-1). After plasma is stored overnight in a tube, these large particles (80 to 500 nm) rise to the top, where they appear as a creamy layer. VLDL contain 90% lipid, the majority being TG, with lesser amounts of cholesterol. When they are present in plasma in increased amounts, VLDL particles are large enough (30 to 80 nm) to impart a cloudy or turbid appearance to plasma. LDL are the major carriers of cholesterol in plasma, and about 50% of their weight is CE and FC. LDL are a heterogeneous group whose size and density vary according to the core content of CE. HDL is often composed of about 50% apolipoprotein and about equal amounts of cholesterol and PL.

Apolipoproteins

Plasma lipoprotein classes are associated with a number of apolipoproteins (Table 12-2). The apolipoproteins have a number of functions; they solubilize lipids in aqueous plasma, allow secretion of chylomicrons and VLDL from intestine and liver, and serve as cofactors for important enzymes in lipoprotein metabolism, and they are responsible for the binding of lipoproteins to specific receptors. The characteristics of the 11 major apolipoproteins and their functions are summarized in Table 12-2. The nucleotide sequences of cDNA for these apolipoproteins have been determined.

LIPID AND LIPOPROTEIN METABOLISM

The transport of plasma lipids by lipoproteins may be divided into an exogenous (dietary) pathway, an endogenous (hepatic) pathway, and reverse cholesterol transport (Fig. 12-1).

Exogenous Lipid Transport

Most of the lipid in the diet is present as neutral fat or TG (75 to 150 g/day). Dietary cholesterol intake is usually about 300 to 400 mg/day but varies from 100 to 600 mg/day. In addition to dietary cholesterol, about 1100 mg of biliary cholesterol is secreted each day from the liver into the intestine (see Fig. 12-1).

In the small intestine, lipids are emulsified by bile salts and hydrolyzed by pancreatic lipases. TG is broken down into FFA and 2-monoglycerides; CE is hydrolyzed into FC and FFA. These lipid components are then absorbed by the intestinal cells. Bile acids are reabsorbed by the intestinal bile acid transporter (IBAT) for return to the liver through the enterohepatic circulation (see Fig. 12-1).

The absorption of cholesterol occurs in the jejunum, through the high-affinity uptake of dietary and biliary cholesterol by the Niemann-Pick C1-like 1 (NPC1L1) protein (see Fig. 12-1). Normally, about half the dietary and biliary cholesterol is absorbed daily. Excessive cholesterol absorption is prevented by the ATP-binding cassette, subfamily G (ABCG5/ ABCG8) transporters, which act together to pump excess cholesterol and plant sterols from the intestine back into the lumen for fecal excretion (see Fig. 12-1).

Inside intestinal cells, monoglyceride is re-esterified into TG and cholesterol is esterified by acyl:cholesterol acyltransferase - (ACAT). Both TG and CE are packaged into chylomicrons, along with apo AI, apo A-IV, and apo B-48. Chylomicrons are secreted into the thoracic duct, from which they enter the peripheral circulation, where they acquire apo C-II and apo E from HDL. Chylomicrons are too large to cross the endothelial barrier, and apo C-II, a cofactor for lipoprotein lipase (LPL), facilitates the hydrolysis of TG near the endothelial lining of blood vessels.

FFA that are released from the hydrolysis of TG are taken up by muscle cells for energy use or by adipose cells for reesterification into TG and storage. As a result, a chylomi cron remnant is produced that is relatively enriched in CE and apo E. This remnant is rapidly taken up by the liver by a process that involves an initial sequestration of remnant particles on hepatic cell surface proteoglycans. Receptor-mediated endocytosis of remnants follows through the interaction of apo E either with the low-density receptor-like protein (LRP, also called the chylomicron remnant receptor)



FIGURE 12-1 Overview of lipoprotein metabolism. Three major pathways of plasma lipoprotein metabolism are shown: transport of dietary (exogenous) fat (left); transport of hepatic (endogenous) fat (center); reverse cholesterol transport (right and center). A detailed description may be found in the text. Sites of action of the six major lipid-altering drugs on exogenous and endogenous pathways of lipoprotein metabolism: (1) inhibition of hydroxymethylglutaryl-coenzyme A (HMG-CoA) reductase by statins; (2) binding of bile acids by sequestrants, interfering with their reabsorption by intestinal bile acid transporter (IBAT); (3) binding of a cholesterol absorption inhibitor to the NPC1L1, decreasing the absorption of dietary and biliary cholesterol; (4) decreased mobilization of free fatty acids (FFA) by nicotinic acid, leading to decreased uptake of FFA by liver and reduced VLDL, IDL, and LDL production; (5) inhibition of TG synthesis by omega-3 fatty acids; (6) upregulation of lipoprotein lipase (LPL) and decreased production of apo C-III, an inhibitor of LPL, by fibric acid derivatives, leading to decreased VLDL-TG. The hepatic cholesterol pool is decreased by the agents at steps 1, 2, and 3, each leading to an upregulation of the LDLR. CETP, cholesteryl ester transfer protein; CMR, chylomicron remnant; HL, hepatic lipase. (From Kwiterovich PO Jr: Clinical and laboratory assessment of cardiovascular risk in children: guidelines for screening, evaluation and treatment. J Clin Lipidol 2:248, 2008. Reproduced with permission.)

or with the LDLR, both on the surface of hepatic parenchymal cells (see Fig. 12-1).

The uptake of dietary and biliary cholesterol is part of a process that regulates the pool of hepatic cholesterol by downregulation of the LDLR and by inhibition of the rate-limiting enzyme of cholesterol biosynthesis, hydroxymethylglutaryl-coenzyme A (HMG-CoA) reductase (see also later). Hepatic cholesterol can be secreted into bile unchanged, converted into bile acids by 7 a -hydroxylase (CYP7A1), or used for lipoprotein synthesis (see Fig. 12-1).

Endogenous Lipid Transport

In the fasting state, most TG in the plasma are carried by VLDL. TG are synthesized in the liver and packaged into VLDL with other lipids and apolipoproteins (see Table 12-1), primarily apo B-100, apo E, apo CI, apo C-II, and apo C-III, and secreted into plasma. The TG in VLDL is subsequently hydrolyzed by LPL, and its cofactor apo C-II, to produce VLDL remnants or IDL. TG can be transferred from VLDL and IDL to HDL and LDL in exchange for CE by the cholesteryl ester transfer protein (CETP; see Fig. 12-1). Compared with VLDL, IDL is relatively enriched

in CE and depleted of TG. Some IDL are taken up directly by the liver, but others are hydrolyzed by hepatic lipase to produce LDL, the final end product of VLDL metabolism (see Fig. 12-1).

The apo B-100 component of the CE-rich LDL binds with high affinity to the LDLR, in either liver or extrahepatic cells (see Fig. 12-1). Bound LDL is internalized by absorptive endocytosis. In lysosomes, apo B-100 is broken down into amino acids, CE hydrolyzed, and FC released. Cholesterol mediates the proteolytic release of a transcription factor, the sterol regulatory element-binding protein (SREBP), from the endoplasmic reticulum. ¹⁷ This effect occurs through the SREBP cleavage-activating protein (SCAP) that is both a sensor of sterols and an escort of SREBP from the endoplasmic reticulum to the Golgi.

The family of transcription factors SREBP regulates the synthesis of cholesterol and fatty acids. SREBP-2 mainly regulates cholesterol synthesis. When hepatocytes are depleted of cholesterol, SCAP transports SREBP from the endoplasmic reticulum to the Golgi, where two proteases, Site-1 protease and Site-2 protease, act in sequence to release the NH $_2$ -terminal domain of SREBP from the membrane. 17

The NH $_2$ -terminal bHLH-Zip domain of SREBP enters the nucleus and binds to a sterol response element in the promoter area of the LDLR and HMG-CoA reductase genes, increasing their transcription. As the cholesterol content of the hepatocyte increases, the SREBP/SCAP complex is not incorporated into the endoplasmic reticulum, SREBP cannot reach the Golgi, and the NH $_2$ -terminal domain of SREBP cannot be released from the membrane for transport into the nucleus, and the transcription of the LDLR and HMG-CoA reductase genes decreases. ¹⁷

This pathway has important clinical implications. For example, excess dietary and biliary cholesterol leads to downregulation of the LDLR and HMG-CoA reductase and an increase in LDL-C. Dietary saturated fat content has a greater effect on LDL-C than dietary cholesterol does. When choles terol is re-esterified by ACAT, SCAP senses a decrease in hepatic cholesterol, leading to upregulation of the LDLR and HMG-CoA reductase genes by SREBP. However, the preferred substrate for ACAT is oleic acid. Thus, excess saturated fatty acids decrease ACAT activity and thereby increase FC, which inhibits the proteolysis and release of SREBP; this downregulates the LDLR and HMG-CoA reductase genes, followed by an increase in LDL-C.

Decrease of dietary cholesterol and saturated fatty acids or of the hepatic cholesterol content with pharmacological agents, such as a cholesterol absorption inhibitor or a bile acid sequestrant (see Fig. 12-1), leads to upregulation of LDLR and HMG-CoA reductase genes and lowers LDL-C. HMG-CoA reductase inhibitors (statins) also reduce hepatic cholesterol content, leading to upregulation of LDLR but without the concomitant increase in HMG-CoA reductase activity (see Fig. 12-1). The concomitant use of a statin with either a cholesterol absorption inhibitor or bile acid sequestrant provides a complementary reduction in hepatic cholesterol and -consequently LDL-C.

When plasma LDL-C exceeds 100 mg/dL, the capacity to process LDL through the LDLR pathway is exceeded. Increased numbers of LDL particles cross the endothelial barrier. LDL is trapped in the vascular wall by proteoglycans and then modified by either oxidation or glycation. Such modified LDL binds to the scavenger receptors CD36 (macro phage scavenger receptor B) and SR-A (scavenger receptor A) and enters macrophages by a low-affinity, LDLR-independent mechanism (see Fig. 12-1).

This pathway is not subject to feedback inhibition of LDLR synthesis by LDL-derived cholesterol. Thus, LDL continues to be taken up in an unregulated fashion, leading to excess deposition of FC and CE in macrophages (see Fig. 12-1). Dys lipidemias that favor an increased uptake of LDL through the scavenger pathway promote the production of foam cells and their associated atherosclerosis and xanthomas.

Pathways of HDL Metabolism and Reverse Cholesterol Transport

Synthesis of HDL

Nascent HDL. Apo A-1 is released as a lipid-free protein from the intestines and liver. Apo A-1 interacts with the ATP-binding cassette transporter 1 (ABCA1) on the basolateral membranes of hepatocytes, enterocytes, and macrophages, acquiring PL and FC to form a more stable nascent HDL particle. ¹⁸

Formation of Larger, Mature HDL Particles. The transition of HDL particles from the disc-shaped nascent HDL to the spherical "mature" HDL requires the esterification of cholesterol to create a hydrophobic core. Lecithin-cholesterol acyl-transferase (LCAT) associates with HDL and catalyzes the transfer of FFA from lecithin to FC, forming CE. Apo AI is a cofactor for this reaction (see Fig. 12-1). CE formed by this reaction constitutes the neutral lipid core of mature spherical -HDL 3; further activity of LCAT provides additional CE for the core of HDL 3, forming the larger HDL 2 particles (see Fig. 12-1). Subsequent addition of cellular cholesterol to the HDL

particle occurs in a number of tissues through the action of another ATP-binding cassette transporter 1 (ABCG1) and the scavenger receptor class B1 (SR-B1), molecules that prefer larger HDL as acceptors. In addition to acquiring lipids from liver and intestines, HDL also acquires lipids from chylomi crons and VLDL in the course of hydrolysis of TG by LPL. During this process, apo AI is transferred from chylomicrons to HDL, and apo C-II and apo E on HDL are transferred to the TG-rich lipoproteins. The TG-rich lipoproteins shed excess PL and cholesterol that are transported to HDL by the phos pholipid transfer protein (PLTP). ¹⁸

Transfer of Lipid Between HDL and the Apo B-Containing Lipoproteins. CE is transferred by CETP from the core of spherical HDL to the apo B-containing lipoproteins in exchange for TG ¹⁹ (see Fig. 12-1). This process depletes CE but enriches TG in HDL particles and has important implications for HDL metabolism. ¹⁸ For example, if the TG in HDL is hydrolyzed by hepatic lipase, a smaller HDL particle is produced that is more avidly removed from plasma by cubilin in the kidney. PLTP is structurally similar to CETP and mediates the transfer of unsaturated fatty acids on PL from the apo B-containing lipoproteins to HDL, contributing to the acquisition of PL by HDL.

Reverse Cholesterol Transport

The CE on the spherical HDL can be transported back to the liver by two mechanisms. CE are transferred from HDL to the apo B-containing lipoproteins by CETP, from which they are taken up by the LDLR (see Fig. 12-1). CE may also be delivered directly to the liver through SR-B1 (also called the HDL receptor). These reactions are part of a process called reverse cholesterol transport.

Once the CE is delivered to the liver and hydrolyzed into FC and FFA, the FC can be excreted directly into bile or converted into bile acids by 7 a -hydroxylase. Both these pathways result in delivery of sterol from peripheral tissues and plasma into intestinal cells, promoting the excretion of sterols into the stool. Reverse cholesterol transport has been postulated to explain, at least in part, the protective effect that HDL and apo AI have against the development of atherosclerosis. Conversely, factors that impede this process, such as a dysfunctional HDL, appear to promote atherosclerosis.

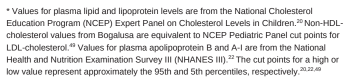
LIPID CHANGES THROUGHOUT CHILDHOOD AND ADOLESCENCE

Clearly, the plasma levels of lipids and lipoproteins result from the influence of a variety of metabolic, genetic, and environmental factors. In the general population, these levels usually follow a Gaussian distribution, often skewed towards higher levels. Percentiles are therefore used to describe such data, with the 50th percentile being the median or average, the 75th to 95th percentiles borderline elevated, and > 95th percentile high or elevated. To define borderline low levels, the 25th to 5th percentiles are used, and < 5th percentile is a low level. ²⁰

Data from the Lipid Research Clinics have often been used to define such percentiles in youth, including the National Cholesterol Education Program (NCEP) Expert Panel on Blood Cholesterol Levels in Childhood in 1992. ²⁰ Similar data are available from such studies as Bogalusa. ²¹ There are also data from the National Health and Nutrition Examination Survey

TABLE 12-3 Acceptable, Borderline, and High Plasma Lipid, Lipoprotein, and Apolipoprotein Concentrations for Children and Adolescents

Category	Acceptable	Borderline	High _*
Total cholesterol	<170	170-199	>200
LDL-cholesterol	<110	110-129	>130
Non-HDL-cholesterol	<123	123-143	>144
Apolipoprotein B	<90	90-109	>110
Triglycerides 0-9 years	<75	75-99	>100
10-19 years	<90	90-129	>130
Category	Acceptable	Borderline	Low⁺
HDL-cholesterol	>45	35-45	<35
Apolipoprotein A-I	>120	110-120	<110



(NHANES) on apo B and apo AI levels 22 and more recent data on TC and LDL-C levels. 23

Human plasma cholesterol levels are lowest during intra uterine life. ²⁴ At birth, the mean (1 SD) plasma levels in normal infants have been reported as follows: TC, 74 mg/dL 11; LDL-C, 31 mg/dL ⁶; HDL-C, 37 mg/dL ⁸; and TG, 37 mg/dL. ¹⁵ The plasma TC and LDL-C levels increase rapidly in the first weeks of life. The kind and source of milk in the infant's diet can significantly influence cholesterol levels. After 2 years of age, the levels of the lipids and lipoproteins become constant up to adolescence. 20 During adolescence, there is a 10% to 20% fall in TC and LDL-C, ²⁵ boys and girls being similarly affected. Also, in the second decade, the TG are higher than they are in the first decade. HDL-C decreases about 10 mg/dL in boys and stays relatively constant in girls. This information has important implications for the timing of lipid and lipoprotein screening and the cut points used because lipid concentrations are age and maturation dependent 26 and appear influenced by gender and race as well, albeit to a small degree. 27

With use of data combined from three major population based prospective cohort studies, TC, LDL-C, HDL-C, and TG variables in childhood and adolescence were classified on the basis of NCEP cut points 20 (Table 12-3), age and gender (not race specific), and NHANES cut points 27 and compared for their ability to predict abnormal levels in adulthood. 28 NCEP cut points (compared with NHANES cut points) were more strongly predictive of high TC, LDL-C, and TG levels in adults but less predictive of low HDL-C.

The continued use of the current NCEP cut points for TC, LDL-C, and TG levels in adolescents appears indicated. Thus, multiple cut points by single age and gender do not appear warranted. NHANES cut points (compared with NCEP cut points) were more strongly predictive of HDL-C in adults. The cut point for HDL-C might be revised upwards, perhaps to 40 mg/dL, to improve the sensitivity of this measurement to predict low HDL-C in adults and to make the cut point congruent with that used in the grown-ups.

Definition of Dyslipidemia

Cut points to define elevated TC, LDL-C, apo B, non-HDL-C, and TG and low HDL-C and apo AI in children and adolescents are found in Table 12-3. Dyslipidemia is present if one or more of these lipid, lipoprotein, or apolipoprotein factors are abnormal.

Significant percentages of children and adolescents have elevated concentrations of lipids and lipoproteins. In the Child and Adolescent Trial for Cardiovascular Health (CATCH), 13% of children in the fourth grade had TC > 200 mg/dL, with 16% of girls and 11% of boys being affected. 29 In the 1988-1994 NHANES, approximately 10% of adolescents had TC > 200 mg/dL. 30 Using data combined from the 1988-2002 NHANES surveys, Jolliffe and Janssen 27 found that the 95th percentile cut points for LDL-C had increased, indicating a shift in the distribution towards higher LDL-C values.

An important epidemiological aspect of cardiovascular risk in childhood is the tracking of lipid and lipoprotein concentrations over time. Tracking indicates the likelihood that children will maintain their percentile ranking over time and has been demonstrated in a number of studies. 21,31,32 In the Mus catine Study, 75% of school-aged children who had TC > 90th percentile at baseline had TC > 200 mg/dL in their early 20s. In the Bogalusa Heart Study, 70% of children with elevated cholesterol continued to have increased cholesterol levels in young adulthood. The Muscatine investigators also found that onset of obesity in adolescence and young adulthood, cigarette smoking, and use of oral contraceptives may have deleterious effects on adult concentrations of lipids and lipoproteins. 31

SCREENING FOR DYSLIPIDEMIA IN YOUTH

Two major approaches have been used to detect dyslipidemia in youth, namely, screening in the general population and in a selected population. The extensive literature related to these two screening approaches has been reviewed in detail. 33 Lipid screening recommendations have recently been updated by the American Academy of Pediatrics. ¹ Traditionally, screening for dyslipidemias in high-risk children has been recommended because of multiple CVD risk factors and a family history of premature CVD or the presence of hypercholesterolemia . LDL-C has been the main focus of diagnosis and treatment.

Less attention has been paid to HDL-C and TG levels. With obesity, type 2 diabetes, and the metabolic syndrome increasing in the younger population, 2,30,34-39 the focus of screening is likely to be expanded to include other factors, such as obesity, low HDL-C, non-HDL-C (TC minus HDL-C), elevated TG, elevated apo B (reflecting increased numbers of LDL particles), glucose intolerance and insulin resistance, and higher blood pressure levels. Both the current and evolving concepts in screening for dyslipidemia in youth are discussed.

Selective Screening

The individualized approach identifies and treats children and adolescents at risk of having high cholesterol levels. Expanded recommendations of the NCEP Expert Panel on Blood Cholesterol Levels in Children and Adolescents 20 from 1992 include performance of selective screening if one of the following conditions is present:

1. A lipoprotein profile in youth whose parents, grand parents, or siblings required coronary artery bypass graft



55 years because of premature CAD.

- 2. A lipoprotein profile in those with a family history of vascular disease, or sudden death before the age of 55 years.
- 3. A TC measurement in those whose parents have high TC levels associated with suboptimal sensitivity, specificity, and predictive TG, low HDL-C, and perhaps Lp(a).
- 4. A lipoprotein profile if the parental or grandparental family result. If one increases the cut point for TC to the 95th percentile usefully expanded to a lipoprotein profile if either obesity (body mass treatment). index >95th per centile) or overweight (body mass index 85th to 94th obesity who do not meet the body mass index criterion.

and Adolescents recommendations 20 from 1992 have been C. 2.45 modified in the most recent American Academy of Pediatrics statement, 1 which recommends screening of children with a family percentile for TC in children is used as a screening cut point, history of premature CVD or high cholesterol or those for whom a about half the individuals who will require treatment as adults family history is unknown or those with other risk factors for CVD are identified by universal lipid screening. In one report, the (eg, obesity, hyper tension, smoking). An optimal program would sensitivity was much lower when screening was performed identify individuals at greatest risk for CVD in adulthood; during adolescence, presumably reflecting the temporary shift of however, there is currently no clinically applicable noninvasive LDL-C to lower values during this period of rapid growth and screening tool available to assess this in children without familial development. 25,46 hypercholesterolemia. Whereas the current targeted approach based on family history assumes that this information is known, elevated TC or LDL-C in children and young adults predicts with adult family members information is often not available.

Universal Screening

Universal lipid screening of all children is controversial . 1,20,33,40 There are a number of advantages and disadvantages of universal screening. What are some of the arguments in favor of universal screening? First, current screening recommendations based on obesity, as part of the screening paradigm for future CVD will family history of CVD or hypercholesterolemia will fail to detect a substantial number (17% to 90%) of children who have elevated lipid levels. 33

undiagnosed heterozygous FH or more marked FCHL, who will require more intensive treatment, usually drug therapy. In a recent meta-analysis of screening for FH in a primary care setting, use of TC detected 88%, 94%, and 96% of cases, with false-positive rates of 0.1%, 0.5%, and 1%, respectively. 41 This approach might usefully will pediatricians and family practitioners handle the detection be combined with a case-finding strategy in relatives of patients with FH. 41

Identification of children and adolescents affected with hypercholesterolemia through universal screening may bring to children, and follow-up be cost-effective? Clearly, national attention their adult relatives who will have greater coronary mortality than relatives of children with normal cholesterol levels. is practiced. ^{2,42} If universal lipid screening is combined with an assessment of obesity and high blood pressure, this can also lead to the detection of additional relatives from families at high risk for CVD. 43

adulthood. Whereas offspring of parents with CVD generally have measured. For selective screening, a lipoprotein profile after higher LDL-C and TG and lower HDL-C in both childhood and young adulthood, 44 the majority of children with

dyslipidemia and multiple risk factors will be missed by 189 selective screening. 33

There are some practical problems with universal screen ing (see later). As well, no longitudinal studies to date are available (and are unlikely ever to be available) to document that starting lipid treatment in childhood decreases adult CVD. 33 One might argue therefore that universal screening seems all the more urgent,

surgery or percutaneous coronary intervention before the age of given the epidemic of obesity and the metabolic syndrome in our youth.

What are potential concerns about universal lipid screening in myocardial infarction, angina pectoris, peripheral or cere bral childhood? The use of TC in childhood to predict TC or LDL-C in young adults, sufficiently high to warrant treatment, is often (> 240 mg/dL). This recommendation might be used fully expanded power of a positive test result. For example, if one uses the 75th to a lipoprotein profile in offspring of parents who have any genetically percentile (about 170 mg/dL) as a lower TC cut point (see Table 12transmissible disorder of lipid metabolism associated with increased 3), the sensitivity (proportion of affected subjects identified) is atherosclerosis risk, involving elevated LDL-C, non-HDL-C, apo B, higher and the specificity (proportion of normal subjects identified as normal) is lower, as is the predictive value of a positive test history is not known and the patient has two or more other risk (about 200 mg/dL), the sensitivity decreases (more children are factors for CAD including obesity (body mass index > 30), missed who are destined to be "affected" as adults), but the hypertension, cigarette smoking, low HDL-C, physical predictive power of a positive test result increases (more test results inactivity, and diabetes mellitus. This recommendation might be above 200 mg/dL correspond to adults who will require

When one uses quantitative traits such as LDL-C values for percentile) is present, regardless of the presence of other non-lipid CVD screening, there is no simple resolution of this problem. ^{2,45} The use risk factors, to identify children and adolescents with abdominal of high LDL-C (> 130 mg/dL) rather than high TC as a cut point improves the sensitivity in those with low HDL-C and the The NCEP Expert Panel on Blood Cholesterol Levels in Children predictive power of a positive test result in those with high HDL-

A number of longitudinal studies 33 found that when the 75th

Another unanswered question is whether the detection of those who are destined to manifest premature CVD. In the Princeton Lipid Research Clinics Prevalence follow-up study of about 30 years, the number (n = 20) of CVD events was small; the sensitivity of childhood LDL-C for prediction of adult CVD was 10.5% and specificity was 81%. 46 Use of family history information does not substantively improve these results. 46 Å combined approach using other CVD risk factors, such as probably improve the detection of those adults more likely to develop premature CVD.

Health care providers need to be aware of the negative impact Universal screening will undoubtedly detect those with of labels on a child. Problems can be created where no problems may have been. Labeling can have a negative impact on a child's self-esteem, potentially predisposing one to a life that may have been different had the labeling never occurred.

> Universal screening raises additional logistical issues. How of many children and adolescents with dyslipidemias? Who will counsel regarding dietary changes, weight loss, and regular exercise habits? Will universal screening, treatment of affected resources will be required to change the way pediatric medicine

What to Measure

CVD risk factors cluster in childhood and persist into A child does not have to be fasting for TC or HDL-C to be

an overnight fast of 10 to 12 hours is measured for screening youth occur well into adulthood. Elevated apo B is the first expression of with a positive family history of premature CVD or dyslipidemia FCHL in adolescents and young adults. 56 or who have obesity, have multiple CVD risk factors, or are suspected of having secondary dyslipidemia. Such a profile includes TC, TG, LDL-C, HDL-C, and non-HDL-C.

Levels of lipoproteins are typically measured and expressed in PRIMARY VERSUS SECONDARY terms of their cholesterol content. LDL-C is calculated from the DYSLIPOPROTEINEMIA Friedewald equation: LDL-C = TC - (HDL-C + TG/5). Total TG in the fasting state divided by 5 is used to estimate the levels of VLDL-C. If the TG is > 400 mg/dL, this formula should not be used and causes must be excluded (Table 12-4). Each child with dyslipidemia a direct LDL-C or apo B should be measured. If the patient is not should undergo routine blood testing to rule out secondary causes, fasting, TC, HDL-C, and non-HDL-C levels can be measured. Non-HDL and apo B are valid in the nonfasting state.

Apo B and apo AI can also be determined by well-12 specific cut points for apo B and apo AI empirically derived from dietary treatment and, if indicated, pharmacotherapy using the the NHANES sample are available, 22 providing cut points that same guidelines as in primary dyslipidemias. can be used to define ele vated apo B and low apo AI (see Table 12-3). Apo B provides an assessment of the total number of apo Bcontaining lipo protein particles. 47

Non-HDL-C is determined by subtracting HDL-C from TC. Non-HDL-C reflects the amount of cholesterol carried by the "atherogenic" apo B-containing lipoproteins – VLDL, IDL, LDL, AND ADOLESCENTS and Lp(a) - and is strongly correlated with apo B. In adults, non-HDL-C is a better independent predictor of CVD than LDL-C is. 47 In children, non-HDL-C is at least as good a predictor as LDL-C of the expression of dyslipidemia in adulthood. ^{2,8,48} Percentiles for non-HDL-C in children are available from Bogalusa 49 (see Table 12-

Advanced lipoprotein testing to determine plasma levels of TABLE 12-4 Causes of Secondary Dyslipoproteinemia VLDL, LDL, and HDL subclasses has been performed in children and adolescents by nuclear magnetic resonance spectroscopy 50-52 Lifestyle or by vertical-spin density-gradient ultra centrifugation 53 in Physical inactivity research studies (see also later). However, cut points derived from Diet rich in fat and saturated fat Alcohol intake these methods for the diagnosis and treatment of dyslipidemia in youth are currently not available.

For universal screening, the simplest approach appears to be the measurement of TC, HDL-C, and non-HDL-C in non- fasting specimens. However, treatment algorithms in pediatric patients are usually focused on fasting LDL-C. TG is usually assessed as part of the dyslipidemic triad and is often ele vated in obesity and the Lipodystrophies metabolic syndrome. 30,34-39,44,54 Thus, in an ideal screening Glycogen storage disease program, follow-up TC, TG, LDL-C, HDL-C, and non-HDL-C in the Acute intermittent porphyria fasted state would be assessed after the initial nonfasting screen.

When to Sample for Dyslipidemia

Cholesterol levels are reasonably consistent after 2 years of age. Cholesterol levels are not routinely measured before the age of 2 years because no formal treatment is recommended for this age group. Ten years of age (range, 9 to 11 years) has been proposed as a good time to obtain a standard lipoprotein profile. 41 Children are older and are able to fast more easily, the values are predictive of Reproduced with permission from Kwiterovich PO: future adult lipoprotein profiles, and puberty has usually not yet started.

Because TC and LDL-C fall 10% to 20% or more during adolescence, 25,27,46 children at risk for familial dyslipidemias should ideally be screened before adolescence, between 2 and 10 years of age. In FH heterozygotes, there is a significant fall in the 1:1 ratio of affected to normal during adolescence. 55 If sampling occurs during adolescence and results are abnormal, levels are likely to be even higher after teenage years. If results during puberty are normal, sampling will need to be repeated towards the end of adolescence (16 years of age for girls and 18 years of age for boys).

The complete phenotypic expression of some disorders, such as FCHL, can be delayed until adulthood, and therefore continued evaluation of subjects from high-risk families with FCHL should

Before a dyslipoproteinemia is considered to be primary, secondary including a fasting blood glucose concentration and kidney, liver, and thyroid function tests. In secondary dyslipidemia, the associated disorder producing the dyslipidemia should be treated standardized immunochemical methods. 22,47 Such measures first in an attempt to normalize lipoprotein levels. If an abnormal might provide additional useful information, particularly in lipid level persists, for example, as it often does with diabetes youth with premature CAD in their parents. 15,16 Age- and gender- mellitus type 1 and nephrotic syndrome, the patient requires

TREATMENT GUIDELINES OF DYSLIPIDEMIA IN CHILDREN

General guidelines for dietary and pharmacological treatment of primary and secondary dyslipidemias in youth are

Endocrine and metabolic

Diabetes mellitus

Metabolic syndrome

Hypopituitarism Hypothyroidism

Pregnancy

Polycystic ovarian syndrome

Renal

Chronic renal failure

Hemolytic-uremic syndrome

Nephrotic syndrome

Hepatic

Biliary atresia

Alagille syndrome

Cirrhosis

Hepatitis

Lipid, apolipoprotein and lipoprotein metabolism. In Kwiterovich PO, editor: The Johns Hopkins textbook of dyslipidemia, China, 2009, Wolters Kluwer, pp 143-156.

Medication

Oral contraceptives

Glucocorticoids

Anabolic steroids 13-cis-Retinoic

Thiazide diuretics Anticonvulsants

Antipsychotics

Estrogen

Testosterone

Immunosuppressive agents

(cyclosporine)

Protease inhibitors

Others

Anorexia nervosa

Cancer survivor

Burns

Idiopathic hypercalcemia

Kawasaki disease Klinefelter syndrome

Progeria (Hutchinson-Gilford

syndrome)

Rheumatoid arthritis

Systemic lupus erythematosus

Werner syndrome

presented first. Specific therapies relevant to each inherited disorder of dyslipidemia are addressed in subsequent sections of this chapter.

Dietary Therapy

The first approach to therapy for children with dyslipidemia is a modified diet containing decreased amounts of total fat, saturated fat, trans -fat, and cholesterol. The intake of complex carbohydrates is increased, and that of simple sugars is decreased. No decrease in total protein is recommended. Adequate calories should be provided to maintain normal growth and development.

The NCEP Pediatric Panel recommends diet treatment after 2 years of age. 20 Recent data from randomized clinical trials in general populations, such as STRIP, indicate that a diet low in total fat, saturated fat, and cholesterol may be instituted safely and effectively at 6 months of age ¹⁰ (see also later).

When to Initiate Dietary Treatment

If the first fasting lipoprotein profile indicates that TC, LDL-C, non-HDL-C, or TG is elevated or the HDL-C is low (see Table 12-3), a repeated profile is obtained at least 3 weeks apart to confirm the first profile. If the dyslipidemia persists (ie, one or more of the lipid persist into childhood and adolescence, when there was no or lipoprotein values remains elevated or the HDL-C is low), relationship of TC to infant feeding. In adults, TC of breast-fed secondary causes of dyslipidemia (see Table 12-4) are ruled out and infants was actually lower than TC of formula-fed infants. Human dietary treatment is initiated. A Step I diet is usually started, and milk remains the "gold standard" for infant feeding. the lipoprotein profile is rechecked in 6 to 8 weeks. If the dyslipidemia persists, a Step II diet is started.

This may not always be available, and the health care provider, soluble fibers 67 such as psyllium 68,69 may also provide an such as the physician or nurse, may need to provide the basis for additional 5% to 10% lowering of LDL-C. The consumption of a soy the diets in Table 12-5 along with printed materials that are protein beverage does not lower LDL-C but may lower VLDL-C available from the American Heart Association. Also, the daily and TG and increase HDL-C. 70,71 estimated calories and recommended servings for grains, fruits, vegetables, and milk or dairy according to age and gender are an omega-3 fatty acid (docosahexaenoic acid, 1.2 g/day) did not available in tabular form from the most recent publication of the lower LDL-C but changed the distribution between the LDL American Academy of Pediatrics. ¹ Assessment of dietary intakes subclasses, with a significant 91% increase in the largest LDL may become possible in the future by use of the web-based subclass and a 48% decrease in the smallest LDL subclass 3.72 Garlic program developed by the National Cancer Institute.

The Step I diet calls for no more than 30% of calories from total 73 fat, < 10% of total calories from saturated fatty acids, < 300 mg/day of cholesterol, and as little trans-fat as possible (only foods with no to be both safe and effective when it is performed under require a reduction in simple sugars and an increase in complex adequacy. carbohydrates. Adequate calories to support growth and Effect of a Low-Fat Diet in Childhood on Future CVD in Adulthood development at a desirable body weight are necessary (this does not appear to be the problem in the United States, however). The Step I diet is evaluated for about 2 to 3 months before the Step II diet is prescribed. The Step II diet entails further reduction of saturated fatty acid intake to < 7% of calories and cholesterol intake from countries with higher CVD and TC levels. ²⁰ Obesity promotes reduction to < 200 mg/day. ²⁰ For a comparison of the Step I and Step II diets, see Table 12-5.

Safety and Efficacy of Dietary Therapy in Infants, Children, and Adolescents

The efficacy and safety of diets low in total fat, saturated fat, and cholesterol to lower LDL-C levels in youth have been demonstrated across the age spectrum of pediatric patients, ² from the age of 7 months to 15 years in STRIP 10,57-59 and from the ages of 8 to 10 years throughout adolescence in the Dietary Intervention Study in Children (DISC). 60 '62 A few studies reported lower intakes of calcium, zinc, vitamin E, and phosphorus on low-fat diets. ² Therefore, whereas

normal growth is achieved and maintained on low-fat diets, adequate intake of these key nutritional elements must be considered.

Owen and colleagues 63 performed a meta-analysis of 37 publications on the effect of breast versus formula feeding on subsequent TC levels in adolescents and adults. Whereas mean TC was higher in breast-fed versus bottle-fed infants, this did not

Nutrient	Step I Diet	Step II Diet
Total fat (total daily calories)	<30%	<30%
Saturated fatty acids	<10%	<7%
	<10%	<10%
Polyunsaturated fatty acids		
Monounsaturated fatty acids	10%-15%	10%-15%
Cholesterol (daily intake)	<300 mg	<200 mg
	50%-60%	50%-60%
Carbohydrates (total daily calories)		
	10%-20%	10%-20%
Protein (total daily calories)		
Total calories	Adequate to achieve and to maintain desirable weight	Adequate to achieve and to maintain desirable weight

The use of margarines (about three servings daily) high in either plant stanol esters 64,65 or plant sterol esters 66 can reduce LDL-C an Both diets require the optimal input of a registered dietitian . additional 10% to 15% when it is added to a low-fat diet. Water-

> Compared with placebo, supplementation of a low-fat diet with extract therapy does not lower LDL-C in hyperlipidemic children.

Overall, a diet low in fat in children with dyslipidemia appears trans -fat on the label are acceptable); given the endemic of supervision. Medical and nutritional support is necessary to overweight and obesity in our society, many children will also reinforce good dietary behaviors and to ensure nutritional -

Prevention of CVD in adulthood by a low-saturated fat, lowcholesterol diet in childhood can be inferred only from epidemiological studies, in which children from countries with a lower prevalence of CVD had lower TC levels than did children insulin resistance in childhood. In that regard, a low-saturated fat dietary program starting at 7 months of age

192 improved insulin sensitivity in 9-year-old healthy children in STRIP 74 and lowered blood pressure in 15-year-olds. 59 This same diet was associated with enhanced endothelial function in boys, but not in girls, mediated in part by the diet-induced reduction in TC. 10 After 10 years of follow-up in STRIP, 10% of the intervention girls were overweight compared with 19% of the control girls, but this significant difference was not seen in boys.

Pharmacological Therapy

Six main classes of lipid-altering drugs are used in children (see Fig. 12-1): HMG-CoA reductase inhibitors (the statins), bile acid sequestrants, cholesterol absorption inhibitors, niacin (nicotinic acid), omega-3 fatty acids (eicosapentae noic acid and docosahexaenoic acid), and fibric acid derivatives.

Institution of Drug Therapy

The primary use of medication in the pediatric population is to lower significantly elevated LDL-C levels, predominantly but not exclusively in children from families with premature CVD or significant dyslipidemia. Drug treatment to lower LDL-C is initiated in children without other CVD risk factors if LDL-C is persistently > 190 mg/dL despite a trial of dietary intervention. Drug treatment is started in children older than 10 years if the postdietary LDLC is > 160 mg/dL and either one or more risk factors for CVD exist, if there is a family history of premature CVD, or if the metabolic syndrome is present. In children with type 1 or 2 diabetes mellitus, pharmacological treatment should be considered when LDL-C is > 130 mg/dL (see also later).

Statins and bile acid sequestrants are the two main classes of pharmaceutical agents currently used in children older than 10 years who have sufficiently elevated LDL-C. Ezetimibe, a cholesterol absorption inhibitor that blocks the absorption of cholesterol and plant sterols through the NPC1L1 protein (see Fig. 12-1), is also effective but is not approved by the Food and Drug Administration (FDA) for use in children, except in rare cases of sitosterolemia or homozygous FH (see later).

Each of these three drug classes reduces hepatic choles terol, leading to the release of the sterol regulatory element binding protein (SREBP) from the cytoplasm into the nucleus, where SREBP binds to the sterol response element in the promoter of the LDLR gene, increasing the number of LDLR and decreasing LDL-C. 17 Because SREBP also upregulates the gene for HMG-CoA reductase, ¹⁷ the bile acid sequestrant and cholesterol absorption inhibitor are both associated with a compensatory increase in cholesterol biosynthesis, limiting their efficacy (see Fig. 12-1). Therefore, both classes of agents might effectively be used in conjunction with statins, which reduce hepatic cholesterol by inhibiting HMG-CoA reductase and decreasing cholesterol biosynthesis.

Niacin is not routinely used in children or adolescents because of its side effects, although some FH homozygotes respond well to niacin (55 to 87 mg/kg/day in divided doses) as a result of a significant reduction of VLDL production, leading to a decreased synthesis of LDL (see Fig. 12-1). In adults, aspirin is used to improve the side effects of niacin. Aspirin is not used in children because of an increased risk for the development of Reye syndrome, so low-dose ibuprofen can be used if necessary to prevent flushing.

The fibrates (48 mg, 135 mg, or 145 mg/day) are also not routinely used in pediatrics, except in the adolescent with a persistently elevated TG level > 500 mg/dL, who may be at increased risk for development of pancreatitis (see also later). Fish oils (2 to 4 g/day) may also be used to treat marked hypertriglyceridemia in children and adolescents by decreasing the biosynthesis of TG (see Fig. 12-1), but the prescription version of omega-3 fatty acids is not yet approved by the FDA for use in children.

Acid Sequestrants Balls

Bile acid sequestrants were the only class of drugs recommended

by the NCEP Pediatric Panel in 1992 for pharmacological lipidlowering therapy because of their record of safety during three decades. 20 In fact, bile acid sequestrants have never been approved by the FDA for use in children. Bile acid sequestrants suffer from significant tolerability issues and provide only a modest LDL-C reduction of about 15%. 76-78 A 17% decrease in LDL-C was reported when cholestyramine (8 g) was used to treat male and female children with heterozygous FH. 77 However, Liacouras and coworkers 78 found that 52 of 63 FH children discontinued cholestyramine treatment after an average of 22 months because of the gritty taste and gastrointestinal complaints.

The second-generation bile acid sequestrant colesevelam (625-mg tablets, three to six per day) has a greater affinity for bile salts and therefore can be used in a lower total dose. In comparison with the first-generation sequestrants, cole-sevelam is associated with less annoying side effects, such as constipation and gritty taste, and interferes less with the absorption of other

In randomized clinical trials, cholestyramine did not affect height velocity. 77 Fat-soluble vitamins were maintained except that the bile acid sequestrant group had significantly lower 25hydroxyvitamin D levels than the placebo group did. One girl had low folate and high homocysteine levels.

HMG-CoA Reductase Inhibitors (Statins)

Statins are widely used in adults to lower TC and LDL-C. A number of randomized, placebo-controlled trials demonstrated the safety and efficacy of statins in male and female FH adolescents. 79-87 A meta-analysis of six of these trials showed high efficacy of statins for LDL-C and apo B lowering and no increase in side effects compared with placebo. 87 Atorvastatin, lovastatin, pravastatin, rosuvastatin, and simv astatin are currently approved by the FDA for use in adolescents with FH. Usual starting doses are atorvastatin, 10 mg/day; lovastatin, 40 mg/day; pravastatin, 40 mg/day; rosuvastatin, 5 mg/day; and simvastatin, 20 mg/day. All except vastatin and rosuvastatin are available generically.

Wiegman and colleagues 85 studied the effect of pravastatin versus placebo on CIMT as a surrogate marker for atherosclerosis (Fig. 12-2). Children aged 8 to 18 years with heterozygous FH were randomized to pravastatin 20 to 40 mg/day (N = 106) or to placebo (N = 108). After 2 years of treatment, those taking pravastatin had a statistically significant mean reduction in LDL-C of 24% versus 0.3% in placebo. Those taking pravastatin had a significant trend towards regression of CIMT compared with those receiving placebo, who had a trend towards progression of CIMT (P < 0.01).

A follow-up study of this Dutch cohort showed that younger age (8 to 10 years) at statin initiation was an independent predictor of the beneficial effect of treatment on CIMT. 88 Early statin therapy also restored endothelial function in children with FH. 89 The American Academy of Pediatrics has recommended that statin treatment be considered in FH children starting at the age of 8 years. ¹ Early intervention with statins appears likely to reduce future atherosclerosis and CVD in those with FH.

Statins may be useful in those adolescents with FCHL or with the metabolic syndrome, who have an LDL-C > 160 mg/dL after diet and weight control, multiple risk factors, or a family history of premature CVD. Statins may also be useful in young women with polycystic ovarian syndrome (see also later), in which there is evidence of increased CIMT, 90 again suggesting that greater attention be paid to management of dyslipidemia and other CVD risk factors early in life.

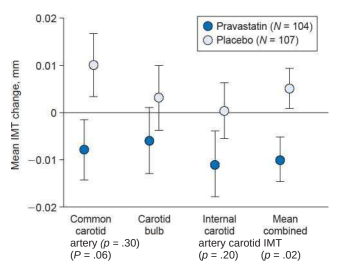


FIGURE 12-2 Mean intima-media thickness (IMT) changes from baseline to 2 years for the different carotid arterial wall segments in the pravastatin (N = 106) and placebo (N = 108) groups of children (aged 8 to 18 years) with heterozygous FH. Compared with the baseline, carotid IMT showed a trend toward regression with pravastatin (mean [SD], -0.010 [0.048] mm; P = 0.049), whereas a trend toward progression was observed in the placebo group (mean [SD], 0.005 [0.044] mm; P = 0.28). The mean (SD) change in IMT between the two groups (0.014 [0.046] mm) was significant (P = 0.02). Pravastatin significantly reduced mean LDL-C levels compared with placebo (24.1% versus 0.3%, respectively; P = 0.001). (From Wiegman A, Hutten BA, deGroot E, et al: Efficacy and safety of statin therapy in children with familial hypercholesterolemia: a randomized controlled trial. JAMA 292:331, 2004. Reproduced with permission.)

Statin Side Effects in Children and Adolescents

Increases in liver function test results up to three times the upper limit of normal were reported in several adolescents treated with higher doses of simvastatin (40 mg/day) 82 and atorvastatin (20 mg/day). 84 In a meta-analysis, 87 the prevalence of elevated alanine aminotransferase in the statin group was 0.66% (3 per 454). Instances of asymptomatic increases (> 10-fold) in creatine kinase (CK), although unusual, have been reported in adolescents receiving statin therapy. 87 No cases of rhabdomyolysis have been reported. 79-87 Adolescents are monitored for elevations in hepatic transaminases and CK concentrations. Liver function test results are checked at each clinic visit two or three times per year. CK is measured at baseline, and measurement is repeated if any myalgia develops.

Special Issues in Young Women

Adult women with FH and CAD may be more responsive than similarly affected men to LDL-C-lowering therapy, as assessed by regression of coronary plaques and tendon xan thomas. 91 The overall favorable safety profile of statin therapy in adult women with CVD is well documented; however, fewer studies have examined the effects of statins in adolescent girls. 82,85,86 Nevertheless, no adverse effects on growth and development or on adrenal and gonadal hormones have been reported. \$2,85,86 One study found a small increase 81 and another study a small decrease 82 in dehydroepiandrosterone sulfate (DHEAS).

Statins are contraindicated during pregnancy because of the potential risk to a developing fetus. Hence, these drugs should be administered to adolescent girls only when they are highly unlikely to conceive. Birth control is mandatory for those who are sexually active.

Because of these concerns, the long-term commitment to therapy, and the fact that CAD often occurs after menopause, some experts believe that statins should not be used to treat adolescent girls. Although the treatment of adolescent patients with FH is indicated, especially of those with a strong family history of premature CAD, additional studies are needed to 193 document the long-term safety of statin therapy and to determine its potential effects on the prevention of atherosclerosis and coronary events.

Metabolic Syndrome—Beyond **Dyslipidemia**

If LDL-C is > 160 mg/dL despite a dietary trial, statin therapy is recommended in patients with the metabolic syndrome. In obese adolescents with metabolic syndrome and LDL-C < 160 mg/dL, the addition of metformin to diet (low in total fat, saturated fat, trans -fat, cholesterol, and simple sugars), exercise, and weight reduction has been shown to be beneficial. 92,93

Treatment of Dyslipidemia Secondary to Other **Diseases**

Diabetes Mellitus

Children with type 1 or type 2 diabetes mellitus are in the highest risk category for the development of manifest CAD early in adult

life. 40 Children with type 1 diabetes and especially type 2 often 12 have a mixed dyslipidemia, the severity of which is related to diabetic control. The American Diabetes Association recommends dietary and other hygienic measures as the first step in the management of these children. However, if the LDL-C is > 160 mg/dL after an adequate trial, the American Diabetes Association panel strongly recommends pharmacological treatment, including the use of statins in adolescents. 94 This recommendation is based on the high risk of CVD in adults with diabetes mellitus type 1 and type 2 and partially on the consistent finding of abnormal CIMT in children with diabetes mellitus type 1.

Nephrotic Syndrome

The dyslipidemia in children with nephrotic syndrome is usually quite marked. The average LDL-C level is close to that found in heterozygotes with FH (Table 12-6). TG can approach 300 mg/dL. The combined elevation of LDL-C and VLDL-C can produce a hypercholesterolemia close to 400 mg/dL. ² Twenty percent of patients with nephrotic syndrome are unresponsive to steroid administration, most cases of which can be attributed to focal segmental glomerulosclerosis. Such individuals with LDL-C > 160 mg/dL may be at an increased risk for the development of atherosclerosis and CVD 1 and may warrant treatment with a

METABOLIC DISORDERS OF DYSLIPIDEMIA IN YOUTH

Disorders Affecting LDL Receptor Activity

There are five disorders expressed in children that result from either mutation in the LDLR or mutations in other genes that affect LDLR activity (Fig. 12-3). Although levels of LDL-C can vary considerably in these five conditions, each disorder manifests early atherosclerosis and premature CVD. These disorders include familial hypercholesterolemia (FH), 95,96 familial ligand defective apo B-100 (FDB), 97 autosomal recessive hypercholesterolemia (ARH), 98,99 sitosterolemia, 100-103 and mutations in proprotein convertase subtilisin/kexin type 9 (PCSK9). 104,105 Each disorder warrants diet and drug therapy in childhood in an attempt to decrease atherosclerosis and subsequent CVD.

Familial Hypercholesterolemia

FH Heterozygotes. FH is the prototype for the diagnosis and treatment of dyslipidemia in children. FH is due to

FH, familial hypercholesterolemia; FCHL, familial combined hyperlipidemia; hyperapoB, hyperapobetalipoproteinemia. Modified from Cortner JA, Coates PM, Gallagher PR: Prevalence and expression of familial combined hyperlipidemia in childhood. J Pediatr 116:514, 1990.



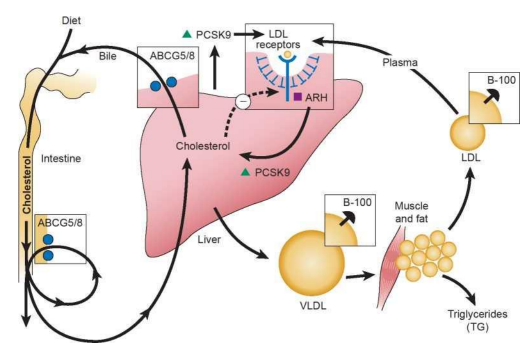


FIGURE 12-3 Schema depicting five inherited disorders of lipoprotein metabolism that are manifested early in life in childhood with marked elevations of LDL, leading to premature atherosclerosis. Apo B, the major apolipoprotein of VLDL and LDL, is necessary for the secretion of VLDL and the uptake of its catabolic product, LDL, by the LDLR. Defects in the structure of apo B-100 (familial ligand defective apo B-100) or in the LDLR (familial hypercholesterolemia) affect the normal binding, internalization, or recycling of the LDLR. Autosomal recessive hypercholesterolemia (ARH) results from a defect in the ARH protein that normally interacts with the cytoplasmic component of the LDLR, allowing the tyrosine phosphorylation and internalization of the LDLR. The PCSK9 is a serine protease that promotes the degradation of the LDLR. Gain-of-function mutations that increase PCSK9 activity decrease LDLR activity. Proposed mechanisms include targeting of the LDLR in the Golgi for degradation in the lysosome, interfering with the recycling of the LDLR after secreted PCSK9 binds to the LDLR at the cell surface, or directing the LDLR to the lysosome to be degraded. The molecular defects responsible for sitosterolemia are caused by mutations in two genes that encode the half-transporters ABCG5 and ABCG8, preventing their normal dual functions of limiting the absorption of cholesterol and plant sterols and promoting their excretion from liver into bile. ABCG5/ABCG8, ATP-binding cassette transporters G5 and G8; PCSK9, proprotein convertase subtilisin-like kexin type 9. (Modified with permission of Goldstein JL. Brown MS: Molecular medicine. The cholesterol guartet. Science 292:1310. 2001.)

mutations in the LDLR gene. Heterozygotes have about a 50% CIMT, 108 decreased brachial endothelial reactivity, 109,110 but rarely reduction in LDLR, whereas homozygotes have little or no LDLR overt CAD. 55 Less than 10% of FH adolescents cent heterozygotes activity. Children with heterozygous FH, an autosomal dominant develop tendon xanthomas. 55 In FH children, the null allele disorder, present at birth 24 and early in life 55 with a twofold to genotype was associated with greater CIMT, higher LDL-C, and threefold elevation in TC and LDL-C 106,107 (see Table 12-6).

When children of an FH parent and a normal parent are compared with receptor-defective mutations. 108 screened, on average half will be affected with FH and half will be normal; in these families, the cut point for LDL-C that minimizes misclassification is about 160 mg/dL. 55 FH affects about 1 in 500 and is due to one of more than 1100 different mutations in the LDLR gene. 95,96 These mutations include insertions, deletions, and missense and nonsense mutations, which can affect the normal synthesis, transport, LDL-binding ability, and clustering (in coated pits) of the LDLR (see Fig. 12-3).

FH heterozygous children and adolescents manifest increased

tendency to attenuate LDL-C lowering with statin therapy

with normals, presumably secondary to the decreased uptake of decreased. 98 IDL by the LDLR, leading to increased IDL and enhanced transfer of TG from IDL to HDL in exchange for CE. About half of untreated chromosome 1 in Sardinian and Lebanese kindred. 98 The ARH adult male heterozygotes and 25% of untreated female protein normally interacts with the cytoplasmic component of the heterozygotes will develop CAD before 50 years of age. 95,96 It is LDLR and other cell surface-oriented molecules, allowing their highly unusual for CAD to occur in adolescents with heterozygous tyrosine phosphorylation. The deficiency of the ARH protein

Treatment of heterozygous FH includes a diet low in cholesterol, total fat, saturated fat, and trans -fat that can usefully manifest a dramatic response to statins alone or in combination be supplemented with plant stanol esters 64 or plant sterol esters 66 and water-soluble fiber. 69 Bile acid sequestrants are safe and moderately effective in FH heterozygotes, 76,77 but long-term compliance is an issue. The dose of the bile acid sequestrant LDL-C level and not to body weight; an adult dose is usually

FH heterozygous children respond well to statin therapy. 79-87 However, the addition of a bile acid sequestrant or cholesterol absorption inhibitor to a statin is often necessary to lower LDL-C to a minimum goal of < 130 mg/dL. Niacin is generally not used to treat FH heterozygous children unless LDL-C is persistently elevated or unusual hypertriglyceridaemia, low HDL-C, or elevated Lp(a) lipoprotein levels are present.

FH Homozygotes. About one in a million children inherit a mutant allele for FH from both parents, leading to LDL-C levels fourfold to eightfold above normal, often causing premature atherosclerosis and death from CVD in the second decade of life. 95,96,107 Atherosclerosis can also affect the aortic valve, leading to severe supravalvular aortic stenosis. Virtually all FH homozygotes have planar xanthomas by the age of 5 years, notably in the webbing of fingers and toes and over the buttocks.

Children and adolescents with the other four disorders affecting LDLR activity (see Fig. 12-3) can also present with planar, tendon, or tuberous xanthomas, as can adolescents with the dominant form of dysbetalipoproteinemia (see later). Secondary disorders of dyslipidemia associated with xantho mas usually have other clinically salient findings to distinguish them from FH homozygotes.

FH homozygotes respond somewhat to high doses of potent statins and to niacin. 111 Because FH homozygotes have markedly diminished LDLR activity, the statins and niacin both work by decreasing hepatic VLDL production, leading to decreased production of LDL-C. Cholesterol absorption inhibitors also lower LDL-C in homozygous FH, especially in combination with a more potent statin. 112 Ultimately, however, FH homozygotes will invariably require regular LDL-C apheresis every 2 weeks to effectively lower LDL-C into a less atherogenic range. 111

Familial Ligand Defective Apo B-100

FDB results from mutations in the gene encoding apo B-100, resulting in an impaired ability of the apo B-100 ligand on LDL-C to bind to the LDLR 95-97 (see Fig. 12-3). For example, the substitution of glutamine for arginine at residue 3500 produces a defective apo B-100 molecule whose binding to LDLR is deficient, leading to decreased clearance of LDL-C from plasma and elevated LDL-C levels.

Heterozygotes for FDB are relatively common (eg, 1 per 1000 in Europeans). 96 About 1 in 20 patients with FDB have tendon xanthomas and appear clinically similar to heterozygous FH patients. Some adult patients with FDB develop premature CAD, but FDB itself is not a common cause of premature CAD. Treatment of FDB is similar to that of heterozygous FH.

Autosomal Recessive Hypercholesterolemia

Children with ARH are clinically similar to those with homo zygous FH, although LDL-C is not usually as elevated (between 350 and 550 mg/dL). 96 In contrast to homozygous FH, both 195 parents of an ARH child usually have a normal lipoprotein profile. Results of functional assays of LDLR in cultured skin

HDL-C is reduced to a certain extent in FH children compared fibroblasts from children with ARH are usually normal or mildly

At least six mutations have been found in the ARH gene on prevents the normal internalization of the LDLR, leading to marked elevations of LDL-C (see Fig. 12-3). Those patients with ARH with a cholesterol absorption inhibitor. 96,98,99

Sitosterolemia

Sitosterolemia (phytosterolemia) is a rare autosomal recessive required to achieve LDL-C < 160 mg/dL is related to the baseline disorder that is expressed in childhood and characterized by markedly elevated (> 30-fold) plasma levels of plant sterols. 96,100,101 This is due to intestinal hyperabsorption and inefficient hepatic excretion of plant sterols and cholesterol. TC and LDL-C can be normal to markedly elevated, depending on the dietary content of cholesterol and plant sterol. Sitosterolemics absorb a higher percentage of dietary cholesterol than normals do, and they secrete less cholesterol into bile, which decreases LDLR activity and in turn increases LDL-C levels 96,100,101 (see Fig. 12-1).

The diagnosis of sitosterolemia is considered and plant sterols are measured in any child or adolescent who has xanthomas despite a disproportionately low LDL-C. Previously undiagnosed adults can mimic FH heterozygotes. Patients with sitosterolemia may develop aortic stenosis, as do those with homozygous FH. 96,100 CVD can present in the first or second decade of life; however, it is usually delayed until early to middle adulthood.

The molecular defects responsible for sitosterolemia are caused by mutations in two genes that encode the half transporters ABCG5 and ABCG8. 96,100,101 These two genes on chromosome 2p are located in a head-to-head orientation. ABCG5 and ABCG8 are expressed exclusively in human liver and intestine, the sites of the two metabolic abnormalities in sitosterolemia (see Fig. 12-3). The dual functions of ABCG5 and ABCG8 are to limit the absorption of cholesterol and plant sterols and to promote their excretion from liver into bile. 96,100,101

Dietary treatment of sitosterolemia is critical, and both cholesterol and plant sterols must be markedly reduced by avoidance of high-fat animal and plant products. Saturated fats are also restricted. Statins are less effective in this disorder because the high sterol content in the liver reduces cholesterol production. ⁹⁶ Bile acid sequestrants are effective, ^{96,102,103} as is ezetimibe. 103 The combination of low-dose cholestyr amine and ezetimibe in a young sitosterolemic girl led to a marked improvement in plasma sterol concentrations, complete regression of xanthomatosis, resolution of carotid bruits, and improvement in her cardiac murmur. 103

Mutations in Proprotein Convertase Subtilisin/Kexin Type 9 (PCSK9)

PCSK9 is a serine protease that facilitates the degradation of the LDLR. 104 Gain-of-function mutations that increase PCSK9 activity decrease LDLR activity, producing a phenotype similar to FH. 104,105 Loss-of-function mutations that decrease PCSK9 activity increase LDLR activity, leading to a lifetime of low LDL-C and a markedly reduced incidence of CVD. 104

The mechanism of action of PCSK9 on LDLR is not completely understood. One possible site of action is in the Golgi, where PCSK9 might target LDLR for degradation in the lysosome. 104 Secreted PCSK9 binds to the LDLR at the cell surface,

leading to the internalization of an LDLR/PCSK9 complex in and the TG levels are normal or moderately elevated. The diagnosis conjunction with ARH (see Fig. 12-3).

PCSK9 may interfere with the recycling of LDLR from the lipoprotein phenotype different from that of the patient. endosome back to the cell surface or direct LDLR to the lyso some to be degraded. It is currently unclear whether PCSK9 cleaves adulthood, 114 it is not unusual to encounter children with FCHL LDLR directly or whether catalytic activity is necessary for either who express different lipoprotein profiles in families with FCHL of these pathways. Patients with hypercholes terolemia and the and premature CAD. In a pediatric lipid clinic, FCHL was three gain-of-function PCSK9 mutation respond well to treatment similar times as prevalent as FH. 106 The mean levels of TC and LDL-C are to that used for FH heterozygotes.

Disorders of VLDL and LDL Overproduction

Phenotypes of dyslipidemia that are due to VLDL overpro duction with FCHL even before the combined dyslipidemia is fully familial combined hyperlipidemia phenotypes are pleiotropic, but the common denominator is of 12 increased numbers of small, dense LDL. Other features can hyperapobetalipoproteinemia, indicating the presence of small, include hypercholesterolemia, hypertriglyceridemia, elevated apo B with normal or borderline LDL-C, low HDL-C, insulin resistance, diabetes mellitus type 2, glucose intolerance, and hypertension. A predilection to CVD is often present. 113

expression of these dyslipidemic phenotypes, indicating that the pathophysiological point of view. No single gene defect in these syndromes has been unequivocally elucidated in humans, and oligogenic factors have been implicated. Treatment of the dyslipidemia (and other aspects) of these phenotypes can successfully start in childhood.

Familial Combined Hyperlipidemia

Goldstein and colleagues 114 described FCHL in families of survivors of myocardial infarction as an autosomal dominant disorder with delayed expression of variable lipoprotein phe notypes: elevated LDL-C level alone (type IIa); elevated LDL-C with hypertriglyceridemia (type IIb); or normal LDL-C with hypertriglyceridemia (type IV).

The prevalence of FCHL is greater than that of FH, occurring in 1 of every 200 to 300 adults. Clinically, it may be difficult to distinguish FCHL from FH if the LDL-C levels are > 200 mg/dL

of FCHL is suspected when a first-degree family member has a

Whereas complete expression of FCHL can be delayed until about 100 mg/dL lower in FCHL children than in children with FH (see Table 12-6). In contrast, mean TG levels are higher in FCHL than in FH children (see Table 12-6).

Total apo B can be elevated in adolescents and young adults (FCHL), expressed. Thus, a child hyperapobetalipoproteinemia, LDL subclass pattern B, familial - hyperapobetalipoproteinemia, as judged by an elevated apo B dyslipidemic hypertension, and syndrome X of Reaven. 113 These level, but normal LDL-C and TG levels (see Table 12-6). The ratio LDL-C В to apo is low dense LDL particles, in contrast to FH, in which the LDL-C/apo B ratio is high, reflecting the underlying large LDL-C particles (see Table 12-6).

Tendon xanthomas are not present in children or adults with Obesity, particularly visceral adiposity, accentuates the FCHL Adult and even adolescent FCHL subjects can develop glucose intolerance, insulin resistance, hypertension, and visceral metabolic syndrome and insulin resistance are intertwined from a obesity. The dyslipidemia of FCHL is often associated with an elevated number of small, dense LDL particles out of proportion to the LDL-C level, a finding that can be evaluated beyond the standard lipid profile by measuring either apo B or the level and size of lipoprotein subclasses by nuclear magnetic resonance spectroscopy. 47

Metabolic Basis of FCHL and Other Small, Dense LDL Syndromes. The abnormal metabolism of FFA in FCHL and other small, dense LDL syndromes may reflect the primary defect in these patients (Fig. 12-4). 113,115 Impaired insulin-mediated suppression of hormone-sensitive lipase in adipocytes leads to an elevation in FFA (see Fig. 12-4). Elevated FFA may drive hepatic overproduction of TG and apo B, leading to a twofold to threefold increased production of

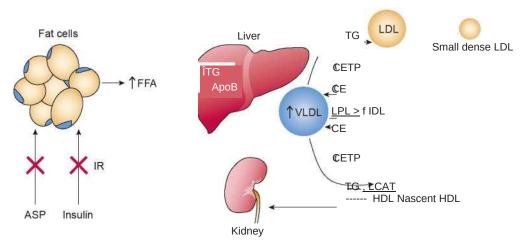


FIGURE 12-4 Mechanisms of the production of the dyslipidemic triad. An increased flux of free fatty acids (FFA) from adipose tissue can result from insulin resistance (IR) or a defect in the acylation stimulatory protein (ASP). Enhanced hepatic uptake of FFA leads to increased production of triglycerides (TG), apolipoprotein (apo B), and VLDL. The TG in VLDL is exchanged for cholesteryl esters (CE) in LDL and HDL by the cholesteryl ester transport protein (CETP), producing CE-depleted LDL and HDL. The TG in the core of LDL and HDL is then hydrolyzed by hepatic lipase (HL), producing both small, dense LDL and smaller HDL. Such HDL is more likely to be excreted by the kidney, resulting in low HDL-C levels. LCAT, lecithin-cholesterol acyltransferase; LPL, lipoprotein lipase. (From Mudd JO, Borlaug BA, Johnston PV, et al: Beyond low-density lipoprotein cholesterol: defining the role of lowdensity lipoprotein heterogeneity in coronary artery disease. J Am Coll Cardiol 50:1735, 2007. Reproduced with permission.)

derived TG-rich lipoproteins.

This paradigm may also result from a cellular defect that prevents the normal effect of acylation stimulatory protein (see Fig. endemic nature in our children. 2,30,34-39 In the past two decades, the 12-4), namely, to stimulate the incorporation of FFA into TG in the prevalence of adolescents with a body mass index > 95th percentile adipocyte. 117 Insulin resistance may also occur in the liver, leading increased by more than 50%. 34 The prevalence of the metabolic to an increase rather than a normal decrease in hepatic syndrome in adolescents increases with the severity of obesity and gluconeogenesis. 118 Finally, FFA and glucose compete as oxidative insulin resistance. 36 Obese adolescents with the metabolic fuel sources in muscle, such that increased concentrations of FFA syndrome also often have the dyslipidemic triad. 34,36 Higher blood inhibit glucose uptake and result in insulin resistance.

Genetic and Molecular Defects

expression of increased particles of small, dense LDL 119 and low HDL-C 120 in FCHL 121,122 and the other small, dense LDL syndromes. 116,121,122 Recently, an orphan G protein-coupled receptor, called C5L2, was reported to bind the acylation stimulatory protein with high affinity and promoted TG synthesis and glucose uptake ¹²³; however, it is not known if C5L2 is defective in hyperapobetalipoproteinemia patients.

Pajukanta and associates 124 mapped the first major gene locus of FCHL to chromosome 1q21-23 and provided strong evidence that the gene underlying the linkage is the upstream transcription factor 1 (USF-1) gene. USF-1 regulates many important genes in lipid metabolism, including the one coding for hepatic lipase, the activity of which is often increased in patients with these syndromes.

Metabolic Syndrome

There is considerable controversy about whether the metabolic syndrome is a discrete clinical entity or simply an aggregation of multiple risk factors for CVD. The characteristics of the metabolic syndrome certainly mirror the pleiotropic phenotypes found in VLDL overproduction. In the Bogalusa Heart Study, metabolic syndrome variables (ie, body mass index, insulin resistance, ratio of TG to HDL-C, and mean arterial pressure) coexisted in terms of Polycystic Ovarian Syndrome childhood and adulthood levels and also in long-term rates of Polycystic ovarian syndrome (PCOS) often presents in adolescence change. 39 Thus, obesity predicted other CVD risk factors both in as menstrual disorders, acne, and hirsutism. 2,90,128 Insulin childhood and in adulthood and was of critical importance in the development of the metabolic syndrome.

Overweight and obesity during childhood, as determined by body mass index cutoff points, are strong predictors of obesity and LDL-C and apo B. After initial diet and weight control, a CAD risk factors in young adulthood. ³⁷ Finally, childhood obesity combination of estrogen and progesterone is used for treatment of and LDL-C are the two strongest predictors of CIMT in young PCOS. 128 adults. 2.7 Clearly, the prevention of obesity should start in

The deleterious effects of obesity are also probably mediated through emerging (nontraditional) risk factors that include atherogenic dyslipidemia. Atherogenic dyslipidemia, also known as the lipid triad, consists of increased numbers of sLDL-P, elevated TG, and low HDL-C; insulin resistance (hyperinsulinemia); a expression and adverse CVD complications in adulthood. proinflammatory state (elevation of serum high-sensitivity Creactive protein); and a prothrombotic state (increased amount of plasminogen activator inhibitor 1).

Specific criteria to screen for multiple CVD risk factors including obesity need to be developed for children. There is no current consensus regarding the definition of the metabolic syndrome in youth. That proposed in children aged 12 to 17 years by Cook and associates ³⁰ from the third NHANES survey is one of several. By use of NHANES data, an adolescent cent was considered to have the metabolic syndrome if three or more of these factors were present: TG > 110 mg/dL; HDL-C < 40 mg/dL; waist patients. $^{2.20}$ The statins have circumference > 90th percentile; fasting glucose concentration > 110 mg/dL; and blood pressure > 90th percentile for age, sex, and height.

One alternative to waist circumference may be a body mass index > 95th percentile for age and gender. Whereas waist circumference and body mass index in children are not routinely

VLDL and the dyslipidemic triad (see Fig. 12-4). 47,113,115,116 Insulin determined, recent data from the Bogalusa Heart Study 125 indicate resistance also interferes with normal upregulation of LPL, leading that both body mass index and waist circumference values, when to decreased lipolysis of TG in VLDL as well as TG in intestinally categorized by a threshold approach, independently predicted CAD risk factors.

> The metabolic syndrome is of paramount interest, given its pressure levels in such adolescents are associated with increased CIMT as a marker for occult atherosclerosis. 35

The presence of the metabolic syndrome in childhood predicts A number of genes (oligogenic effect) ² may influence the the adult metabolic syndrome, diabetes mellitus type 2, and the development of CVD two to three decades later. 126,127 The finding of acanthosis nigricans reflects the insulin resistance that is often present. In addition, biomarkers for increased risk of atherosclerotic disease, such as high-sensitivity C-reactive protein and adiponectin, were increased and decreased, respectively, in obese children. 36

Measurement of lipoprotein subclasses by nuclear magnetic resonance spectroscopy (despite the fact that this is not yet recommended for routine clinical practice) provides some insights into the relationship of lipoprotein heterogeneity and obesity and other components of the metabolic syndrome. 2 TG, insulin, and relative weight in children from an unselected population were positively associated with the size of VLDL and negatively associated with the size of LDL particles. 50 Large VLDL but not small VLDL was notably higher in white than in black children across quintiles of waist circumference. 51 The presence of fatty liver obese adolescents was associated with a pronounced dyslipidemic profile characterized by large VLDL, small dense LDL, and decreased concentration of large HDL particles. 52 This pro-atherogenic phenotype was strongly related to the intrahepatic lipid content.

resistance, considered one of the underlying causes of PCOS, has increased substantially in the past decade, putting more adolescent girls at risk for PCOS and its complications, including elevated

Only about one in three specialists consider metformin appropriate treatment in teenagers with PCOS; however, in obese teenagers with PCOS, almost 70% would use metfor min. 128 Adults with PCOS appear to be at high risk for CVD. 90 Increased CIMT occurs in young adults with PCOS, 90 indicating that early diagnosis and treatment of PCOS in adolescence may prevent its full

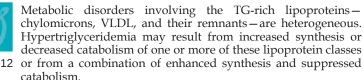
Treatment of Disorders of VLDL Overproduction

A low-fat diet reduces the burden of postprandial chylomi crons and the atherogenic chylomicron remnants (see Fig. 12-1). Reduction to ideal body weight improves insulin sensitivity and decreases overproduction of VLDL. Regular aerobic exercise is of paramount importance. Two classes of drugs, fibric acids and niacin, lower TG and increase HDL-C in adults and may also convert small, dense LDL particles into larger LDL particles. 113 However, fibrates and niacin are not ordinarily used in pediatric

198 the most effective in lowering LDL-C and the total number of atherogenic, small, dense LDL particles. 113

In adolescents with FCHL or with the metabolic syndrome, drug treatment with a single agent, most often a statin, is reserved for those with a more marked elevation of LDL-C > 160 mg/dL (see also earlier). Cholestyramine has also been used to treat pediatric patients with FCHL who have elevated LDL-C. 78 Metformin has been used to treat obese hyperinsulinemic adolescents with the metabolic syndrome. 92,93 Metformin can enhance insulin sensitivity and reduce fasting blood glucose concentration, insulin, lipids, FFA, and leptin. 92,93

Familial Metabolic Disorders of Triglyceride-Rich Lipoproteins



Most hypertriglyceridemia in children and adolescents is due to VLDL overproduction, often accompanied by obesity or overweight and other components of the metabolic syndrome (see earlier). The focus here is on inherited disorders of marked hypertriglyceridemia. The most serious complication of marked hypertriglyceridemia is pancreatitis. Some but not all families with hypertriglyceridemia will ultimately manifest CVD in middle age.

Endogenous Hypertriglyceridemia

Familial Hypertriglyceridemia. In some children, TC and LDL-C levels are normal but VLDL-C and TG are elevated (type IV lipoprotein pattern). Familial hypertriglyceridemia (FHTG) can be distinguished from FCHL by demonstrating that affected parents and siblings of the proband have normal LDL-C and apo B levels (see Table 12-3). In contrast, relative to FCHL, these levels are borderline high or elevated (see Table 12-5).

Whereas VLDL particles in FHTG and FCHL are both TG enriched, VLDL and apo B are not being overproduced in FHTG as they are in FCHL. HDL-C is low or normal in FHTG. Adults with FHTG often manifest glucose intolerance, obesity, hyperuricemia, and peripheral vascular disease. The disorder may be inherited as an autosomal dominant trait with reduced penetrance in childhood; for example, only one in five children younger than 20 years born to an affected parent expresses FHTG.

Exogenous Hypertriglyceridemia

Two classic inherited disorders result from absent or markedly deficient activity of LPL: a defect in LPL and a defect in apo C-II, the cofactor for LPL (see Fig. 12-1). 129 A newer unique defect of chylomicronemia has recently been described, a deficiency of glycosylphosphatidylinositol-anchored high-density lipoprotein-binding protein 1 (GPIHBP1) associated with chylomicronemia. 130 GPIHBP is an endothelial cell protein that normally binds LPL and chylomicrons. A homozygous mutation (Q115P) in GPIHBP1 eliminates the normal binding of GPIHBP

Defective or Missing Lipoprotein Lipase. Defective LPL produces a profound hypertriglyceridemia (as high as 10,000 mg/dL) because of the massive increases in chylomi crons and the inability to clear dietary fat. 129 Marked hyper cholesterolemia (eg, 300 to 1000 mg/dL) is also present secondary to hyperchylomicronemia, with a ratio of TG to TC of at least 5 and usually 10. VLDL-C is normal, and HDL-C and LDL-C are low. LPL activity is absent in plasma and adipose tissue of patients with this disorder (also called type I hyperlipoproteinemia).

The half-life of chylomicrons is prolonged about sixfold. The plasma levels of apo C-II, the cofactor for LPL, are normal. The

diagnosis requires measurement of plasma lipolytic activity after intravenous injection of heparin (60 units/kg), which releases the membrane-bound lipases into the blood stream (post heparin lipolytic activity [PHLA]).

This disorder is usually manifested early in the first year of life. Creamy blood is often noted after a blood draw or in a hematocrit tube after a fingerstick. Abdominal pain is a common symptom that is manifested as colic during infancy or as an acute abdominal discomfort later in childhood. Other clinical features may include eruptive xanthomas, hepato-splenomegaly, and lipemia retinalis.

Premature atherosclerosis does not occur in LPL deficiency because chylomicrons are too large to enter the vascular wall and are therefore not atherogenic. LPL deficiency is a rare recessive trait. Obligate heterozygous parents of affected children are often consanguineous and have normal lipid levels or a moderate dyslipidemia. More than 80 mutations in the LPL gene have been reported. 129

Defect in Apolipoprotein C-II. When apo C-II, the cofactor for LPL, is deficient, hypertriglyceridemia can range from 800 to almost 10,000 mg/dL; the lipoprotein pattern may be type I, or the elevated chylomicrons may be accompanied by ele vated VLDL (type V). 1.129 TC is normal or increased (151 to 980 mg/dL). LDL-C and HDL-C are low (< 5th percentile). PHLA is absent or very low. Plasma apo C-II is present in only trace amounts.

If apo C-II is added to plasma in vitro or by blood or plasma transfusion in vivo, normal PHLA is restored. This disorder is a rare, autosomal recessive trait. Pancreatitis, which usually manifests itself in adulthood, is the major clinical problem. Pancreatitis developed in one homozygous child by 6 years of age. Abnormalities of the apo C-II gene are caused by either small deletions or by splice-site mutations. 129

Defect in GPIHBP1. Sixty patients with severe hypertriglyceridemia were screened for mutations in GPIHBP1, an endothelial cell protein that binds LPL and chylomicrons. Mice lacking GPIHBP1 had been found to develop TG levels above 2000 mg/dL even on a low-fat chow diet. 130 The ability of GPIHBP1-expressing CHO cells to bind LPL and chylomicrons suggested that this protein may function as a platform for lipolysis on endothelial cells in heart, skeletal muscle, and adipose tissue.

One 33-year-old man with life-long chylomicronemia was found to be homozygous for a GPIHBP1 (Q115P) mutation. He had failure to thrive as a child but no history of pancreatitis . After a PHLA test, he had low levels of LPL at 10% of normal, the cause of which requires further study. He responded to a low-fat diet, with the TG level falling from 3366 mg/dL to 744 mg/dL.

Treatment of Profound Exogenous Hypertriglyceridemia

Strict fat restriction to 10 to 15 g/day is required. Intake of linoleic acid as ~1% of total calories must be maintained. Medium-chain TG (MCT), which are absorbed directly through the portal vein and therefore do not require mobilization of chylomicrons from the intestine into plasma, can be added to the diet as 15% of calories to make the diet more palatable. Infants with a type I pattern may take a formula high in MCT.

A subset of LPL-deficient children with unique, possibly posttranscriptional genetic defects respond to therapy with MCT oil or omega-3 fatty acids by normalizing fasting TG levels. Thus, a therapeutic trial with MCT oil should be considered in all children presenting with the familial chylomicronemia syndrome. 2,131 A dramatic response to antioxidant therapy was reported in a small number of patients with LPL deficiency for whom dietary measures

failed and in whom frequent severe episodes of pancreatitis developed.

Endogenous and Exogenous Hypertriglyceridemia

Type V Hyperlipoproteinemia

Patients with marked hypertriglyceridemia may also present with increased chylomicrons and VLDL (type V lipoprotein phenotype). Common clinical findings include pancreatitis, eruptive xanthomas, retinal lipemia, abnormal glucose tolerance , and insulin resistance. Because LPL and apo C-II are not deficient, lipolysis of TG on both chylomicrons and VLDL occurs, albeit at a reduced rate. This leads to the production of both chylomicron remnants and VLDL remnants; thus, both peripheral and coronary atherosclerosis can develop in type V

Although type V is usually expressed in young adulthood, this phenotype has been reported in several preadolescent children. Affected relatives may have either type V or type IV lipoprotein phenotypes. Autosomal dominant inheritance has been described in several large families.

Elevated VLDL may result from increased synthesis, decreased clearance, or a combination of both, but the fundamental defect in many of these patients has remained elusive. In two unrelated male and female patients with type V, Cao and colleagues 132 found two different heterozygous frameshift mutations in *CAV1* that encode caveolin 1. Mice with a deleted CAV1 develop adipocyte abnormalities and insulin resistance. Thus, the genomic DNA from patients with atypical lipodystrophy and hypertriglyceridemia (with no mutations in any known lipodystrophy gene) was screened for defects in the coding regions of human CAV1.

Dysbetalipoproteinemia (Type III Hyperlipoproteinemia)

Patients with dysbetalipoproteinemia present with elevated TC and TG levels, often above 300 mg/dL. The hyperlipidemia is caused by the accumulation of chylomicron remnants derived from intestinal lipoproteins and VLDL remnants derived from hepatic lipoproteins.

Dysbetalipoproteinemia is conventionally defined by a cholesterol-rich VLDL that has \$ rather than pre- \$ electropho retic mobility. ^{2.133} Ultracentrifugation of plasma, without density adjustment (density < 1.006 g/mL), is required for the diagnosis to be made. The ratio of VLDL-C to plasma total TG is usually > 0.3 (normal, 0.15 to 0.25). Such "beta" VLDL reflects the accumulation of both chylomicron and VLDL remnants.

This recessive defect involves a polymorphic genetic locus that specifies the structure of apo E. ¹³³ Apo E binds to receptors on the surface of hepatocytes, promoting uptake of both chylomicron and VLDL remnants. Human apo E exists as three major isoforms (apo E2, apo E3, and apo E4), each of which is specified by an independent allele at the locus for the apo E gene. The most common allele is apo E3. The necessary but insufficient cause of type III hyperlipoproteinemia is the presence of two copies of apo E2, which differ from the most common isoform of apo E (apo E3) by a single amino acid substitution, which is associated with the recessive form of dysbetalipoproteinemia. 133

One in 100 people is homozygous for the apo E2 allele. Most patients with dysbetalipoproteinemia have apo E2 homo zygotes; because the prevalence of the disorder is low at 1:10,000, other modifying factors, such as an overproduction of VLDL in the liver (seen in FCHL) or hormonal and environmental conditions (hypothyroidism, low-estrogen state, obesity, or diabetes mellitus), are necessary for complete clinical expression. This recessive form of dysbetalipoproteinemia has a delayed penetrance. The dominant form of dysbetalipoproteinemia can be expressed in childhood and **199** does not require the presence of modifying factors. However, clinical features of the disease are often delayed until adulthood.

In both the recessive and dominant forms of dysbetalipoproteinemia, affected patients often develop xanthomas, particularly yellowish deposits in the palmar creases (xanthoma striatum palmare) and tuberous and tuberoeruptive xantho mas over the elbows, knees, and buttocks. Tendon xanthomas are much less frequent. Premature atherosclerosis of the coronary, carotid, abdominal, and femoral arteries is prevalent. Hyperuricemia and glucose intolerance occur in up to half of the patients with this syndrome.

Treatment of the Combined Exogenous and **Endogenous Triglyceride Disorders**

Treatment starts with a diet low in total and saturated fat, transfat, cholesterol, and simple carbohydrates but higher in complex carbohydrates and protein (see also earlier) and weight management with reduction to ideal body mass. If TG level remains > 500 mg/dL, drug therapy, including fibrates, niacin, fish oils (omega-3), and statins (see Fig. 12-1), can be considered starting in adolescence, particularly in patients who have a strong family history of premature CVD or a history of eruptive xanthomas or pancreatitis.

Familial Disorders of HDL Metabolism

In most instances, low HDL-C occurs secondary to VLDL overproduction (see Fig. 12-3) and is expressed as a component of the dyslipidemic triad. There are, however, primary HDL-C disorders that present with low HDL-C levels and CVD: familial hypoalphalipoproteinemia 134,135; homozygous gene deletions or nonsense mutations in apo AI 18; missense mutations in apo AI 136,137; more than 100 common and rare variants in ABCA1, including the prototype, Tangier disease 138; and LCAT deficiency. 139

One disorder, CETP deficiency, often presents as high HDL-C, but whether this is associated with increased or reduced risk for CVD is not resolved. Inhibition of CETP with torcetrapib increased large HDL but was associated with an increased total and CVD mortality. 140

Hypoalphalipoproteinemia

Hypoalphalipoproteinemia is defined as low HDL-C (< 5th percentile or < 35 mg/dL; see Table 12-3) in the presence of otherwise normal lipids and number of sLDL-P. Adults with this syndrome exhibit an increased prevalence of CVD; however, such patients do not manifest the clinical findings typical of severe forms of HDL deficiency, such as planar xanthomas, corneal clouding, and peripheral neuropathy. 18

hypoalphalipoproteinemia, although prevalent than the rare recessive disorders, including deficiencies in HDL, is relatively uncommon. 134,135 In some families, hypoalphalipoproteinemia behaves like an autosomal dominant trait and can be associated with genetic variants. 18

Apolipoprotein Al Variants

A number of apo AI variants have been described and are attributed to specific amino acid substitutions. 2,18,136,137 For example, apo AI Milano results from a mutation in the apo AI gene at codon 173, changing arginine to cysteine. 137 Heterozygous carriers for autosomal codominant traits often have low HDL-C but are usually asymptomatic with regard to premature CVD. Intravenous injection of recombinant apo AI Milano weekly for 6 weeks appeared to reduce coronary atherosclerosis in adults with acute coronary syndromes. 137



Tangier disease 138 is a rare metabolic disorder, originally described by Donald Fredrickson in two young girls from Tangier Island in the Chesapeake Bay, 141 in which HDL is both abnormal and present in severely reduced concentration. Tangier homozygotes have plasma apo AI levels < 3% of normal. Immunochemically detectable apo AI is synthesized by intestinal

cells but is rapidly degraded in plasma.

HDL in Tangier disease (termed HDL r) is markedly abnormal; a chylomicron-like lipoprotein particle is present in the density range of HDL on a normal high-fat diet. These abnormal lipoproteins are rich in CE and are likely to be sequestered by the reticuloendothelial cells in Tangier disease. These large, flattened, lucent particles, 100 nm in diameter, disappear when a low-fat diet is consumed. These observations suggest that HDL is necessary for normal metabolism of chylomicrons.

The compositions and amounts of the other lipoproteins are also abnormal. TC is decreased, with normal or elevated TG. The 12 lipoprotein abnormalities are accompanied by a striking deposition of CE in different tissues. The major clinical manifestations reflect the lipid storage and include enlarged orange-yellow tonsils, splenomegaly, and a relapse of peripheral neuropathy. Mild hepatomegaly, lymphade nopathy, and corneal infiltration (in adulthood) may also appear. Foam cells can be demonstrated on biopsy of the skin, bone marrow, peripheral nerves, or rectum.

The apo AI gene is normal in Tangier disease. The basic defect resides in mutations in ABCA1. 138 Thus, ABCA1 was discovered by virtue of studies in Tangier patients from three laboratories. ABCA1 normally mediates the efflux of cellular cholesterol onto the nascent HDL particle for transport to the liver (see Fig. 12-1).

Lecithin-Cholesterol Acyltransferase Deficiency and

LCAT is an enzyme bound to HDL-C (a -LCAT) and to a lesser extent to VLDL/LDL-C (§ -LCAT) in the plasma. 139 LCAT catalyzes the formation of CE in lipoproteins. The two familial forms of LCAT deficiency are termed familial LCAT deficiency (complete LCAT deficiency) and fish eye disease (partial LCAT deficiency). Both familial LCAT deficiency and fish eye disease are autosomal recessive disorders.

In classic LCAT deficiency, both a -LCAT and \$ -LCAT activities are absent, resulting in a markedly reduced plasma cholesterol esterification rate, a low plasma cholesteryl ester content, and an abnormal lipoprotein profile with very low HDL-C. Clinical findings include glomerulosclerosis, normochromic anemia, corneal opacities (detectable in childhood), and premature atherosclerosis. Specific defects in the LCAT gene, including stop codons and amino acid substitutions, have been elucidated in several kindred with classic LCAT deficiency. 139

Fish eye disease is a phenotypically distinct syndrome of LCAT deficiency in which most but not all patients appear to have a selective defect in a -LCAT activity, which is accompanied by dense corneal opacities; low HDL-C is present, but premature atherosclerosis does not develop. Several molecular defects have been described in the LCAT gene of patients with fish eye disease. An interesting mutation, LCAT(300-del), has led to the postulate that the heterogeneity in the phenotypic syndromes of LCAT deficiency may be related to the residual amounts of total plasma LCAT activity.

Treatment. In Tangier disease, a low-fat diet diminishes the abnormal lipoprotein species that are believed to be remnants of abnormal chylomicron metabolism. The large LDL-C species found in LCAT deficiency is also thought to be a remnant of abnormal chylomicron metabolism. Its disappear ance on a lowfat diet may have a beneficial effect because large LDL-C may be involved in the pathogenesis of renal disease. Patients with other syndromes associated with deficiencies of HDL-C and premature atherosclerosis are also treated with a diet significantly reduced in total fat, saturated fat, trans -fat, and

cholesterol.

Hyperalphalipoproteinemia

In distinct contrast to hypoalphalipoproteinemia, some children have very high HDL-C levels (> 95th percentile), termed hyperalphalipoproteinemia. TC concentration is often ele vated as a result of elevated HDL-C levels; LDL-C concentration is usually normal, and TG levels are normal or low. Hyperalphalipoproteinemia is often associated with longevity and a decreased risk of CVD. Hyperalphalipoproteinemia can be due to a defect in CETP. Lower SR-B1 levels and two SNPs in the SR-B1 gene (see Fig. 12-1) were found in subjects with hyperalphalipoproteinemia. 142

Decreased activity of endothelial lipase due to loss-offunction mutations in LIPG constitutes a novel genetic basis of familial hyperalphalipoproteinemia. 143 Several kindred with hyperalphalipoproteinemia have been described in which affected members have defects in hepatic lipase (see Fig. 12-1). Mutations in hepatic lipase are associated with elevated HDL-C levels in the paradoxical presence of hypertriglyceridemia . Few data are available on the expression of hepatic lipase defects in children.

Elevated Levels of Lp(a) Lipoprotein

Lp(a) lipoprotein is a very large molecule (M _r 3 x 10 ⁶) found in the density range of 1,050 to 1,080 g/mL. ¹⁴⁴ Its lipid composition is similar to LDL, but Lp(a) contains two proteins, apo B-100 and a large glycoprotein called apo(a). Apo(a) is attached to apo B-100 through a disulfide bond. Apo(a) is homologous to plasminogen with a yet uncertain biological function and has a variable number of repeats of the kringle 4 region, which are under genetic control. An inverse relationship exists between the size of apo(a) and the level of Lp(a). 144

At present, most laboratories that measure Lp(a) use commercially available antibodies. Elevated Lp(a) appears to be inherited and is often strongly associated with premature CVD in some families. An Lp(a) level is usually elevated in children with strokes and should be measured as part of the workup in every child. ¹ Niacin is the only lipid-altering drug that reduces Lp(a). It is currently unknown if treatment of elevated Lp(a) prevents development of future or recurrent CVD; however, current recommendations are to aggressively lower LDL-C when the Lp(a) is high.

Deficiencies in Apo B-Containing Lipoproteins

The discussion here is limited to those genetic conditions of low LDL-C levels and reduced CAD. For the rarer primary or secondary disorders of deficiencies of apo B-containing lipo proteins that present in early childhood, the reader is referred elsewhere. 145

Hypobetalipoproteinemia

The phenotype of primary hypobetalipoproteinemia is characterized by very low levels of LDL-C, usually defined as the lower 5th percentile of a normal distribution. Plasma TC concentration is low; VLDL-C and TG levels are low or normal.

Familial Hypobetalipoproteinemia. hypobetalipoproteinemia is inherited as an autosomal dominant. A relatively large number of mutations in APOB causing hypobetalipoproteinemia have been described. 146 Almost all



of the mutations are either nonsense mutations or frameshift mutations that create a premature stop codon and a truncated apo B-100. The endogenous VLDL system (see Fig. 12-1) is defective because the apo B-100 product of the normal allele is produced at approximately 25% of normal, and the truncated apo B is cleared too rapidly from plasma. ¹⁴⁶ This results in decreased production of apo B-100 and VLDL and low LDL-C (see Fig. 12-1).

Familial hypobetalipoproteinemia has also been linked to a susceptibility locus on the chromosome 3p21 and in some families is linked neither to *APOB* nor to chromosome 3p21. ¹⁴⁶ Familial hypobetalipoproteinemia patients are usually asymptomatic, the prevalence of CVD is low, and often longevity is found. Those with a defect in apo B can have increased hepatic fat of about threefold.

Loss-of-Function Mutations in PCSK9. The phenotype of hypobetalipoproteinemia is also found in those with a loss-of-function mutation in the PCSK9 gene. ¹⁴⁷ In this case, the low LDL results not from decreased production of LDL but from enhanced LDLR activity due to the decreased PCSK9 function ^{147,148} (see Fig. 12-3). Patients with this cause of familial hypobetalipoproteinemia also have a considerable life long reduction in CVD, ^{147,148} presumably starting at birth or early in life.

CONCLUSION

Early lesions of atherosclerosis begin in childhood and are related to antecedent CVD risk factors. Environmental factors such as diet, obesity, and exercise and certain inherited dyslipidemias influence the progression of such lesions. Detection of youth at risk for atherosclerosis includes an integrated evaluation of these predisposing factors. Treatment starts with a diet low in saturated fat, *trans* -fat, and cholesterol. Such intervention can be supplemented with water-soluble fiber, plant stanols or plant sterols, weight control, and exercise.

Drug therapy, especially with HMG-CoA reductase inhibitors or bile acid sequestrants, can be considered in those with a positive family history of premature CVD and LDL-C > 160 mg/dL, after a period of dietary and hygienic measures. Candidates for drug therapy often include those with FH, FCHL, the metabolic syndrome, polycystic ovarian syndrome, type 1 or type 2 diabetes, and the nephrotic syndrome. Such dietary and drug therapy appears to be safe and effective.

The early identification and treatment of children with CVD risk factors and dyslipidemia are likely to retard the atherosclerotic process. Optimal identification and treatment of high-risk youth either from the general population or from families with premature CVD will require an integrated universal screening, evaluation, treatment, and follow-up program.

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CHAPTER 13

The Role of High-Density Lipoprotein Cholesterol in the Development of Atherosclerotic Cardiovascular Disease

Puneet Gandotra and Michael Miller

KEY POINTS

- HDL-C is inversely correlated with coronary heart disease.
- In addition to its pivotal role in reverse cholesterol transport, HDL possesses antiinflammatory, fibrinolytic, and antioxidant properties.
- Levels of HDL-C inadequately characterize HDL functionality.
- Therapies that raise HDL-C levels are associated with reduced atherosclerotic progression.
- Clinical outcome studies are evaluating whether raising of HDL-C levels independently reduces coronary heart disease risk.

EPIDEMIOLOGY AND PATHOPHYSIOLOGY CONSIDERATIONS

Epidemiology

During the past several decades, numerous observational studies have strongly supported high-density lipoprotein cholesterol (HDL-C) as an independent risk factor for coronary heart disease (CHD). In the United States, the Framingham Heart Study provided the strongest examples of this relationship. ¹ As demonstrated in Figure 13-1, the risk of CHD was highest at the lowest levels of HDL-C in the Framingham Heart Study, even when lowdensity lipoprotein cholesterol (LDL-C) levels were not elevated (ie, < 100 mg/dL). Conversely, high levels of HDL-C conferred relative cardioprotection compared with lower HDL-C levels, even when they were accompanied by high LDL-C levels (ie, > 220

Inverse relationships between HDL-C and CHD have also been demonstrated outside of the United States as exemplified in Troms0, Norway, where a threefold greater risk of future CHD was conferred by low HDL-C independently of any other variables. 2 In Europe, the western Prospective Cardiovascular Munster (PROCAM) study observed more than 25,000 men and women without symptomatic CHD at baseline (1979-1991). Among the most prominent findings was that low HDL-C, defined as < 35 mg/dL, conferred a 2.5-fold increase of incident CHD in the absence of elevated total cholesterol (< 200 mg/dL) and a 5-fold increase at higher total cholesterol levels. 3

Other prospective studies confirmed HDL-C to be inversely correlated with incident myocardial infarction even after adjustment for other risk factors, such as age, smoking, blood pressure, weight, and diabetes mellitus. ⁴ If low HDL-C predicted CHD events, might there be a useful metric to gauge clinical effects related to raising HDL-C levels? In fact, before clinical outcome studies addressing the effect of raising HDL-C on CHD events (see

later), observational data from the Multiple Risk Factor Trial, ⁵ the Lipid Research Clinics ⁶ follow-up trial, the placebo arm of the Coronary Primary Prevention Trial, ⁷ and the Physicians' Health Study ⁸ suggested that an HDL-C increment of 1 mg/dL would translate into an approximate 3% reduction in CHD risk.

Taken together, observational data from previous decades (1970s-1990s), with few exceptions, 9-11 paint a vividly convincing picture in favor of HDL-C as an independent risk factor for CHD. Similarly, clinical trials assessing the effect of lipid-altering therapy on outcomes found baseline measurements of low HDL-C to be predictive of both initial and recurrent CHD events. 12 Moreover, patients with low HDL-C assigned to placebo therapy demonstrated increased atherosclerotic progression and possessed the highest CHD risk in clinical trials evaluated in statin therapy. 13

Pathophysiology of High-Density Lipoprotein

In addition to reverse cholesterol transport, HDL possesses antioxidant, anti-inflammatory, and antifibrinolytic properties, all believed to contribute to its putative athero-protective role. However, the extent to which these properties translate into clinical improvement vis-a-vis CHD outcomes awaits the results of ongoing clinical trials (see later).

The physiological pathway underlying reverse cholesterol transport is shown in Figure 13-2. ¹⁴ Originating as an HDL precursor, lipid-depleted and hepatically or intestinally derived apolipoprotein (apo) AI receives phospholipids and free cholesterol hydrolysis of triglyceride-rich lipoproteins. The relatively lipid-poor pre- \$ (based on electrophoretic mobility) HDL interacts with the ATP transport protein ABCA1 to shuttle free cholesterol and phospholipids from peripheral cells (eg, macro phages). Cholesterol is sequestered into the HDL core through esterification by

lecithin-cholesterol acyltransferase (LCAT) to form small spherical HDL $_3$ (or a $_2$, a $_3$ HDL) 15 particles with approximately two molecules of apo AI and a small percentage of particles containing hepatically derived apo A-II. 16

Additional contributions of cholesterol mediated by ABCG1 and ABCG4 (and, to a minor extent, the scavenger receptor B1 [SR-B1]) result in HDL maturation and larger HDL 2 particles (or a 1) that contain three or four molecules of apo AI. Whereas apo AI plays an important role in stabilizing HDL, activating LCAT and promoting reverse cholesterol transport, the role of apo A-II in this process is less clear. ¹⁶ The fate of cholesteryl esters contained within HDL includes transfer to lower density lipoproteins (eg, LDL, VLDL) in exchange for triglyceride, a process mediated by the cholesteryl ester transfer protein (CETP).

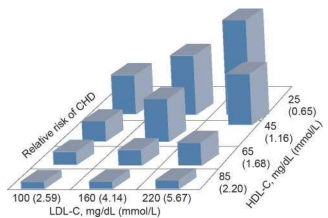


FIGURE 13-1 Risk of coronary heart disease (CHD) by HDL-C and LDL-C levels from the Framingham Heart Study. (From Harper CR, Jacobson TA: New perspectives on the management of low levels of high-density lipoprotein cholesterol. Arch Intern Med 159:1049, 1999.)

The cholesteryl ester transferred to very-low-density lipoprotein (VLDL) is converted to LDL and taken up by LDL receptors for hepatobiliary delivery and excretion. Alternatively, HDL may deliver cholesteryl ester to steroidogenic tissues (liver, adrenal, testis, ovaries) by binding to a SR-B1 protein, when it may serve as a precursor for hormone and gonadal steroid production. In cases of CETP inhibition (see later), apo Eenriched HDL may also deliver hepatic cholesterol by LDL receptor-related mechanisms. Other less established contributors to reverse cholesterol transport include apo AI uptake by high-affinity hepatic beta-chain ATP syn thase receptors ¹⁷ and by the renal proximal tubule endocytic receptor cubilin. ¹⁸

In addition to its role in reverse cholesterol transport, HDL possesses other atheroprotective properties (Fig. 13-3). Specifically , HDL downregulates the expression of interstitial vascular cell adhesion molecule 1 (ICAM-1) and vascular cell adhesion molecule 1 (VCAM-1) in vascular endothelial cells. This limits transendothelial migration of monocytes and macrophage conversion, thereby reducing a proinflammatory milieu. In addition, the HDL-associated enzymes paraoxonase 1 (PON1) and platelet-activating factor acetylhydrolase (PAF-AH) inhibit LDL oxidation. ^{19,20} The inverse association between PON1 activity and CHD ²¹ validates the potential clinical relevance of non-reverse cholesterol transport characteristics of HDL.

After the discovery that apo AI is a prostacyclin-stabilizing factor, ²² attention has been drawn to the HDL-associated phospholipid sphingosine 1-phosphate (S1P) because of an array of biological effects on vascular endothelial and smooth muscle cells. For example, HDL-associated S1P improves arterial tone by upregulation of endothelial nitric oxide synthesis . Moreover, in cellular studies and animal models, administration of HDL-associated S1P reduced infarct size and protected endothelial cells against apoptosis. ²³

HDL has also been shown to reduce platelet activation by upregulating prostacyclin and nitric oxide production

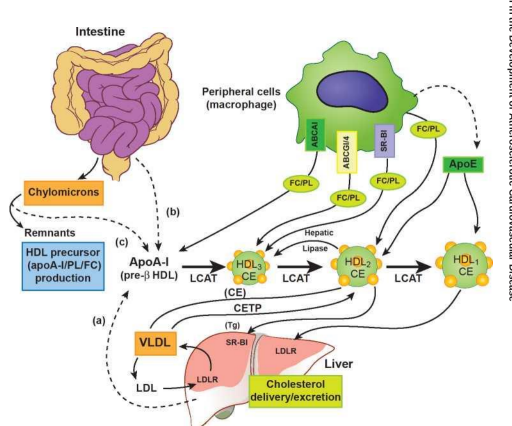


FIGURE 13-2 Role of HDL in the redistribution of lipids from cells with excess cholesterol (e.g., macrophages) to cells requiring cholesterol or to the liver for excretion. CE, cholesteryl ester; CETP, cholesteryl ester transfer protein; FC, free cholesterol; LCAT, lecithin-cholesterol acyltransferase; LDLR, LDL receptor; PL, phospholipid; SR-B1, scavenger receptor B1; Tg triglyceride. (From Mahley RW, Huang Y, Weisgraber KH: Putting cholesterol in its place: apo E and reverse cholesterol transport. J Clin Invest 116:1226, 2006.)

Beyond reverse cholesterol transport and other putative atheroprotective properties is the recent identification of serine protease inhibitors and complement-modulating proteins associated with HDL (Fig. 13-4) that may protect against proteolysis and plaque rupture. ²⁵ Although in its early development stages, HDL proteomics is likely to advance our understanding of the diverse roles that HDL plays in vascular biology and atherothrombosis.

HIGH-DENSITY GENETICS **LIPOPROTEIN**

High-Density Lipoprotein Deficiency States and Coronary Heart Disease Risk

To date, chromosomal aberrations in the APOA1/C3/A4/A5 gene complex or mutations in ABCA1, LCAT, and APOAI have been implicated in HDL deficiency states (eg, HDL-C < 10 mg/dL). Yet, in the absence of other CHD risk factors,

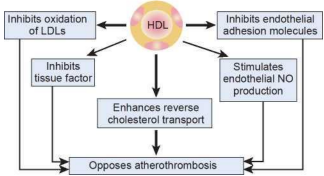


FIGURE 13-3 Other potential atheroprotective roles attributable to HDL. NO, nitric oxide. (From Barter P: Metabolic abnormalities: high-density lipoproteins. Endocrinol Metab Clin North Am 33:393, 2004.)

premature CHD, especially before the age of 40 years, has not been reported with hereditary HDL deficiency. This is in marked contrast to familial hypercholesterolemia, in which, regardless of risk factor status, homozygotes commonly develop CHD in childhood or adolescence and heterozygotes may manifest CHD before the age of 30 years. ²⁶

The lack of premature CHD in the absence of other risk factors (eg, smoking, diabetes) suggests that there are other mechanisms enabling cholesterol efflux as the initial step in reverse cholesterol transport. For example, in cases of ABCA1 deficiency, cholesterol efflux occurs through SR-B1, passive diffusion, or upregulation of sterol 27-hydroxylase (CYP27A1) and caveolin 1 (Fig. 13-5). 27 Similarly, in the absence of apo AI, other apolipoproteins such as macrophage-derived apo E may contribute to reverse cholesterol transport. 28

A third possibility is that from a teleological standpoint, HDL was designed to remove only excess cholesterol to maintain cellular lipid homeostasis. If this is correct, "isolated" low HDL-C as defined by physiological levels of LDL-C and triglyceride (eg, less than 70 mg/dL and 100 mg/dL, respectively) without traditional CHD risk factors would not be associated with increased risk of premature CHD.

In fact, as noted before, HDL-C deficiency as a consequence of monogenic abnormalities has rarely if ever been associated with premature CHD in the absence of accompanying CHD risk factors, most notably cigarette smoking. ²⁹ Rather, the increased CHD risk associated with low levels of HDL-C may in large part be ascribed to associated metabolic disturbances (eg, visceral adiposity, insulin resistance) that upregulate proinflammatory signaling pathways and raise overall atherothrombotic risk. This is in striking contrast to familial hypercholesterolemia, in which, in the pre-statin era, premature CHD occurred in most affected subjects, regardless of risk factor status.

In contrast, elevated levels of HDL-C (ie, > 60 mg/dL) have been viewed as a negative CHD risk factor. 30 Yet, the association of heritable high HDL-C (> 60 mg/dL) with reduced CHD risk remains as debatable as isolated low HDL-C and premature CHD. For example, variation in the HDL regulatory gene CETP is associated with high HDL-C and is especially prevalent in Japan, where approximately 50% of high HDL-C is a consequence of genetic CETP deficiency. ³¹ Despite several case reports suggesting reduced CHD risk (Table 13-1), data are inconclusive as to whether intrinsic cardioprotection is afforded as a consequence of CETP inhibition.

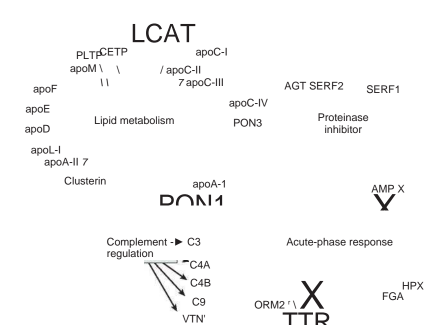


FIGURE 13-4 Global view of biologic processes and molecular functions of HDL proteins. (From Vaisar T, Pennathur S, Green PS, et al: Shotgun proteomics implicates protease inhibition and complement activation in the antiinflammatory properties of HDL. J Clin Invest 117:746, 2007.)

TABLE 13—1 Selected Cases of Cholesteryl Ester Transfer Protein

Deficiency and Evidence of Atherogenicity*

		HDL-C		
Mutation	Location	(mg/dL)	Atherogenicity	Reference
Int14A	Japan	>70	Т	139
D442G	United States	<60	Т	140
		>60	4	
R37X	Sweden	208	4	141
R268X	Canada	70	4	142
IVS7+1	Netherlands	80	T4	143
Q87X/Q165X	Greece	194	4	144

*Defined by history of coronary heart disease or noninvasive evaluation (carotid intimamedia thickness, computed tomographic angiography).

TABLE 13-2	Novel Candidate Genes Associated with HDL-C			
genes	Chromosomes HDL s		s	Reference
GALNT2	1	Т		145, 146
PCYT1	3	Т		147
ELOV2	6	4		148
VNN1	6	Т		149
BMP1	8	4		150
MMAB/MVK	12	4		151

To address the potential benefit of CETP inhibition, a multicenter trial is currently evaluating whether the CETP inhibitor dalcetrapib reduces CHD outcomes (see later). In addition to *CETP*, variation in hepatic lipase (*LIPC*) and endothelial lipase (*LIPG*) has also been associated with high levels of HDL-C. However, *LIPC* variants have not been associated with cardioprotection, ³² and it remains unclear whether the loss-of-function *LIPG* variant Asn396Ser affects atherogenicity. ³³

In addition to loss-of-function variants that despite being rare in the general population (< 1%) produce a significant phenotypic effect (eg, very low or high HDL-C), genome wide association studies investigate informative loci that may also contribute to HDL-C levels. In addition to known genes that regulate HDL metabolism and function, novel HDL genes have recently been uncovered through genome-wide association studies, ³⁴ a sample of which is represented in Table 13-2.

The potential mechanism of action of these encoded proteins on HDL metabolism is outlined in Figure 13-6. Although the impact on CHD risk has yet to be defined, the addition of genome-wide association studies to rare variant identification is likely to provide new insights related to HDL metabolism and overall atherothrombotic risk.

High-Density Lipoprotein Functionality and Classification

One of the quagmires relating HDL to CHD risk assessment is how best to assess and to represent HDL functionality. Although the level of HDL-C is most commonly used to gauge

CHD risk, it inadequately characterizes the HDL proteome in terms of inherent metabolic complexities and effectiveness of reverse cholesterol transport. This in part reflects the inherent

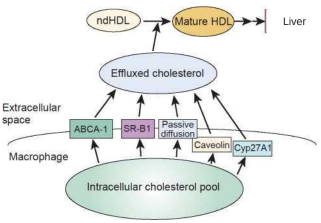


FIGURE 13-5 Cholesterol efflux may occur by at least five independent routes, including ATP-binding cassette transporter A1 (ABCA1), scavenger receptor B1 (SR-B1), caveolin, Cyp27A1, and passive diffusion. (From Ohashi R, Mu H, Wang X, et al: Reverse cholesterol transport and cholesterol efflux in atherosclerosis. QJM 98:845, 2005.)

difficulty in accurately quantifying the contribution of macrophage-derived fecal cholesterol, a small measure of reverse cholesterol transport. Nevertheless, the recent development of macrophage-specific reverse cholesterol transport assays in murine models ³⁵ may hold promise as a future diagnostic tool if results are reproducible in humans.

In addition to cholesterol efflux and reverse cholesterol transport, other putative measures of HDL-mediated atheroprotection are currently under exploration. They include evaluation of reconstituted HDL, apo AI Milano and mimetic compounds on indices of inflammation, hemostasis, thrombosis, and endothelial function. ³⁵

Although HDL-C levels have traditionally been used as a surrogate for reverse cholesterol transport, experimental evidence favors additional measurements that may contribute to CHD risk stratification. For example, in the setting of insulin resistance or in hypertriglyceridemic states (Fig. 13-7), free fatty acids are mobilized from adipocytes to drive hepatic VLDL production. The enhanced synthesis of triglyceride-rich lipoproteins leads to upregulation of CETP, resulting in greater exchange of triglyceride and cholesteryl ester between VLDL and HDL. Hypertriglyceridemic HDL particles exhibit reduced efficiency of cholesterol efflux ³⁶ and are subsequently hydrolyzed by hepatic lipase to produce small, dense cholesterol-depleted HDL particles.

Apo AI is catabolized by cubilin receptors in the proximal renal tubule, ³⁷ and higher apo AI fractional catabolic rates account for the reduced HDL-C levels found in postprandial states ³⁸ as well as in obese, insulin resistance, and diabetic states. This in part reflects upregulation of CETP in triglyceride-enriched apo B-100 containing lipoproteins (most notably, VLDL), thereby allowing greater exchange of triglyceride-cholesteryl ester with HDL. The triglyceride-enriched HDL serves as an excellent substrate for hepatic lipase, resulting in small, dense, apo AI-depleted HDL particles. ³⁸

Whereas greater apo AI stability might suggest a more resounding cardioprotective effect for larger HDL particles, observational studies have been inconsistent. For example, several studies have found HDL 2 to be associated with lower CHD risk, ³⁹ whereas others, including the Physicians' Health Study ⁴⁰ and EPIC-Norfolk prospective population study, ⁴¹ failed to find significant differences between HDL particle





PC biosynthesis

FIGURE 13-6 Novel genes and their proposed effects on HDL metabolism. SIRT1 activates the liver X receptor (LXR), thereby leading to upregulation of ABCA1 and cholesterol efflux. Mevalonate kinase (MVK) catalyzes an early step in cholesterol biosynthesis, and the MMAB (methylmalonic aciduria) gene encodes a protein involved in cholesterol degradation. GALNT2 regulates glycosylation of proteins such as LCAT. VNN1 inhibits lipid peroxidation, a cardioprotective property of HDL. PCYT1 encodes a protein regulating the production of phosphatidylcholine, an integral component of HDL. BMP1 converts a precursor to mature form of apo AI. (From Holleboom AG, Vergeer M, Hovingh GK, et al: The value of HDL genetics. Curr Opin Lipidol 19:385, 2008.)

FC biosynthesis

. WVK (MMAB?)\

Lipid peroxidation

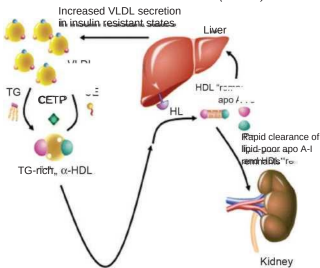


FIGURE 13-7 Mechanism of decreasing HDL-C and apo A-I in insulin-resistant and other hypertriglyceridemic states. Insulin-resistant states are associated with an increase in VLDL production and postprandial chylomicronemia. Cholesteryl ester transfer protein (CETP)-mediated exchange of HDL cholesteryl ester (CE) with triglyceride (TG) results in CE depletion and TG enrichment of HDL particles. Hepatic lipase (HL), which is also increased in insulin-resistant states, modifies TG-rich HDL, releasing lipid-poor apo A-I and forming HDL "remnant" particles. Lipid-poor apo A-I can be either recycled to form mature spherical HDL particles or filtered by the renal glomerulus and then degraded by proximal renal tubular cells. HDL remnants may also bind to putative receptors in liver or kidney that mediate HDL uptake, internalization, and degradation. (From Rashid S, Patterson BW, Lewis GF: Thematic review series: patient-oriented research. What have we learned about HDL metabolism from kinetics studies in humans? J Lipid Res 47:1631, 2006.

remnants," and lipid-poor Al and apo A-II

size and risk of incident CHD after adjustment for traditional risk factors and other covariates.

A final lingering concern is the potential conversion of HDL to a proinflammatory form (Fig. 13-8). 42 For example, in the setting of an acute coronary syndrome (ACS), the enzyme myeloperoxidase released from leukocytes may bind to and oxidatively modify apo AI. In addition to reducing the effectiveness of apo AI in reverse cholesterol transport, 43 the associated nitration and chlorination of apo AI convert HDL to a proinflammatory particle that may be selectively incorporated in human atheroma. 44

Moreover, displacement of apo AI by the acute-phase reactant serum amyloid A and associated reductions in LCAT and the HDL antioxidants PON1 and PAF-AH eliminate many of the cardioprotective properties of HDL. Statins may offset some of the adverse effects associated with proinflammatory HDL particles after ACS. 45

THERAPIES THAT AFFECT HIGH-DENSITY LIPOPROTEIN

Lifestyle Changes Leading to Increased Levels of HDL-C (Table 13-3)

Diet and Weight Loss

The International Diabetes Federation and National Choles terol Education Program consider abdominal adiposity and low HDL-C part of the metabolic syndrome. Weight loss, especially when it is accompanied by aerobic activity (see also later), may have a significant impact on raising HDL-C levels. ⁴⁶ In the absence of weight loss, however, increasing the proportion of carbohydrates at the expense of fat reduces both HDL-C and LDL-C levels. ⁴⁷ Moreover, the active process of weight loss is commonly associated with transient reductions in HDL-C, especially when a low-fat diet is prescribed, ⁴⁸ because of reductions in apo AI production. ⁴⁹

However, once weight loss has been achieved and body weight stabilized, a meta-analysis of 70 diet studies found minimal increases approximating an increase of 1.6 mg/dL in HDL-C for every 10 pounds of weight lost. ⁵⁰ Finally, high intake of marine-based omega-3 fatty acids has been associated with higher HDL-C in some populations, ^{51,52} although the overall net increase is very modest. Specifically, fish oil consumption by diet or capsule form has been associated with a 3% increase in HDL-C in subjects without hypertriglyceridemia at baseline (ie, < 177 mg/dL) compared with no increase in HDL-C among subjects with higher triglyceride levels. ⁵²

Exercise

Aerobic exercise has been shown to increase HDL-C levels on average by 5% to 10%, with the increase mostly related to the frequency and intensity of the exercise. ⁵³ Increased HDL levels with exercise are associated with upregulation of lipo protein lipase activity. ⁵⁴ Overall, 10% to 20% increases in HDL-C are observed if at least 1200 kcal are expended weekly. ⁵⁵

Conditioned athletes and endurance runners often have HDL-C levels that are 30% to 50% higher than those of sed entary subjects, ⁵⁶ probably reflecting the combination of enhanced lipoprotein lipase activity and increased production of pre-Ş HDL particles, which may facilitate reverse cholesterol transport. ⁵⁷ Smaller increases (5% to 10%) are observed in subjects who have baseline low levels of HDL-C accompanied by elevated triglycerides and visceral adiposity. ⁵⁸On the other hand, isolated low HDL-C is difficult to effectively raise even after extended endurance training. ⁵⁹

Smoking Cessation

Overall, cigarette smoking impairs LCAT activity ⁶⁰ and reduces levels of HDL-C. In women, 1 pack per day smokers evidenced HDL-C levels that were 10 mg/dL lower compared with those of nonsmokers, whereas an approximate difference of 3 mg/dL in HDL-C favoring nonsmokers was observed in men. ⁶¹ In contrast, a meta-analysis of 27 studies found

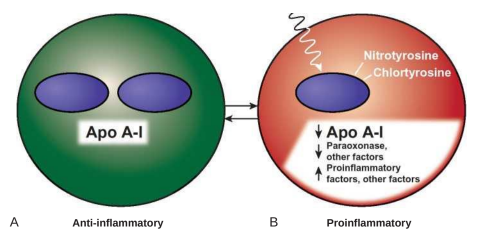


FIGURE 13-8 Model of bidirectional conversion of HDL from anti-inflammatory (A) to proinflammatory (B) and the role of myeloperoxidase in catalyzing oxidative modification of HDL, rendering it unable to effect ABCA1-mediated cholesterol transport. In association with these changes, apolipoprotein A-I (apo A-I) and paraoxonase levels decrease, whereas proinflammatory factors such as lipid oxidation products increase. (From Ansell BJ, Fonarow GC, Fogelman AM: The paradox of dysfunctional high-density lipoprotein. Curr Opin Lipidol 18:427, 2007.)

TABLE 13—3 Effect of Lifestyle Modification on HDL-C Levels			
Lifestyle Component	Percentage Increase in HDL-C		
Diet and lifestyle	0-5		
Weight loss	5-15		
Omega-3 fatty acids	< 5		
Exercise	5-20		
Cigarette cessation	5-10		
Alcohol consumption	5-15		

significant increases in HDL-C levels after smoking cessation. ⁶²

In a study performed by Moffatt, ⁶³ participants who stopped smoking for 60 days saw an average increase of 12.5 mg/dL in HDL-C; reinitiation of cigarettes resulted in reversion to precession levels.

Alcohol

Alcohol inhibits hepatic lipase, thereby raising both HDL-C and its HDL $_2$ subfraction. 64 A dose-response relationship exists; 1 ounce of alcohol consumed daily is associated with up to a 15% increase in HDL-C. 65 It has been suggested that raising HDL levels represents approximately half of the CHD benefit attributable to alcohol use. 66

The type of alcoholic drink does not appear to be as important as the quantity. 67 HDL-C increases due to alcohol intake may be more noteworthy in sedentary subjects compared with those who exercise on a regular basis. 68 Caution should be exercised in patients with low HDL-C who have elevated triglycerides (ie, > 200 mg/dL) as alcohol may substantially raise these levels as well.

Pharmacological Therapy

In addition to lifestyle recommendations, medications have been used to raise levels of HDL-C. It remains to be established whether and to what extent raising HDL-C in and of itself reduces CHD risk. We review current and investigation agents affecting HDL metabolism as well as ongoing clinical trials that it is hoped will provide insights into the virtue of raising HDL-C

or improving its functionality.

Niacin

Nicotinic acid, or niacin (vitamin B $_3$), is currently the most potent HDL-C-raising medication in the United States, with increases ranging between 20% and 35%. 69 Of the three for mulations, immediate-release (IR; three times daily), slow-release (SR; twice daily), and extended-release (ER; once daily), the IR form raises HDL-C to the greatest extent, followed by ER and SR formulations. For example, at doses of 1000 mg, the IR formulation raises HDL-C 25%, compared with 15% to 20% with ER and 10% with SR formulations. 70,71 At doses of 1500 to 2000 mg daily, observed increases range from 25% to 35% (IR) versus 20% to 30% (ER) and 10% to 20% (SR).

Niacin reduces HDL catabolism by inhibiting hepatic apo AI removal. ⁷²Niacin also inhibits adipose hormone sensitive lipase, reducing free fatty acid flow to hepatocytes, thereby decreasing VLDL production and triglyceride output. ⁷³Based on the inverse association between triglycerides and HDL-C, for every reduction of 50 mg/dL in triglyc erides, HDL-C levels rise by approximately 1.6 mg/dL in addition to apo AI-mediated effects. ⁷⁴Preliminary data for a combination of niacin and statin therapy found increases in HDL ³ levels of apo J and phospholipid transfer protein, raising the specter of improved reverse cholesterol transport, although the direct contribution by niacin (and statin) treatment remains to be determined. ⁷⁵

In the Coronary Drug Project (CDP), 3 g of IR niacin daily was associated with reduction in nonfatal myocardial infarction after 5 years and a total mortality benefit of 11% compared with placebo. ⁷⁶ Coronary arteriography and studies evaluating carotid intima-media thickness have demonstrated that the addition of niacin to statin-based therapy is associated with reduced progression and in some cases regression of atherosclerotic disease. ⁷⁷

For example, in the Arterial Biology for the Investigation of the Treatment Effects of Reducing Cholesterol 3 (ARBITER 3), the addition of niacin to statin therapy resulted in decreased progression and mild regression of carotid intima media thickness at 12 and 24 months, respectively. ⁷⁸ More over, the HDL Atherosclerosis Treatment Study (HATS) found that the combination of niacin and statin decreased atherosclerosis by angiography as well as decreased clinical events compared with placebo. ⁷⁹ In this study, despite a low sample size, those randomized to the combination (compared



with placebo) had a significant 90% lower rate of recurrent cardiac events, noteworthy given that most statin studies have shown maximum event reductions on the order of 30% to 40%. set the stage for the clinical outcome trial Atherothrombosis Intervention in Metabolic Syndrome with Low HDL/High Triglycerides and Impact on Global Health Outcomes (AIM-HIGH) designed to evaluate whether the combination of niacin and statin (with or without ezetimibe) is clinically superior to a predominant LDL-C-lowering regimen alone. 80

The most common side effect of niacin therapy is the prostaglandin D 2 -mediated cutaneous reaction that includes flushing and, to a lesser extent, urticaria. Pretreatment with 325 mg of chewable or nonenteric aspirin may limit or abort this reaction. Whereas the intensity of this side effect is reduced with food and ER or SR formulations and often eases with continued administration, instances of renewed flushing may occur even with long-term administration as a result of insufficient food intake, overconsumption of alcohol, overexposure to heat, and aerobic activity.

In this regard, a selective prostaglandin D 2 receptor 1 antagonist (laropiprant) that is associated with reduced flushing is currently being studied in a clinical outcome trial (see also later). 81 Other less frequent niacin-based side effects include dyspepsia, gout, acanthosis nigricans, toxic amblyopia, and elevation of plasma glucose concentration. However, studies suggest that diabetic subjects receiving up to 3 g IR or 2 g ER niacin had relatively modest (5% to 7%) but clinically insignificant increases in fasting glucose levels. 82-84 In fact, a CDP post hoc analysis of diabetic and metabolic syndrome patients found that niacin therapy improved CHD outcomes. 76

The most disconcerting side effect, hepatic toxicity, has not been encountered with the ER regimen. 83,84 Nevertheless, the results of the AIM-HIGH trial and the Heart Protection Study 2-Treatment of HDL to Reduce the Incidence of Vascular Events (HPS2-THRIVE) comparing the combination of nicotinic acid/laropiprant and statin therapy versus statin based therapy alone 85 will in large part dictate the future of this therapy in patients with vascular disease.

Fibrates

Fibrate therapy generally results in a 10% to 25% increase in HDL-C levels. 86 Fibrates are synthetic agonists of peroxisome proliferator-activated receptor a (PPAR a), which stimulates expression of the hepatic apolipoprotein AI gene 87 and modulates the transcription of other genes involved in reverse cholesterol transport, including SR-B1 and ABCA1. 88 However, the relative increase in HDL-C is largely driven by the associated reductions in triglyceride levels, the primary lipid-mediated effect of fibrates. 89 In primary prevention, the Helsinki Heart Study of 4081 asymptomatic middle-aged men with elevation of non HDL-C (> 200 mg/dL) showed the fibrate gemfibrozil to be associated with a 34% overall reduction in incident CHD. 90 However, lower median HDL-C at baseline (ie, < 42 mg/dL) was associated with improved CHD outcomes despite more modest raising of HDL-C compared with higher baseline HDL-C levels.

Moreover, in the secondary prevention Veterans Affairs HDL Intervention Trial (VA-HIT), recurrent CHD events were reduced by 11% with every increase of 5 mg/mL in HDL-C when patients were treated with gemfibrozil. 92 In this study, the HDL subfraction HDL 3 was correlated to a greater degree than total HDL-C levels with CHD. Moreover, gemfibrozil treatment resulted in a 10% increase in HDL particle number that correlated with a 29% reduction in CHD risk. 93

Statins

Although statins exert a more modest effect on HDL-C levels (5% to 15% increase), they are especially effective in patients with low HDL-C, in whom they may attenuate the elevated risk associated with reduced levels (see Fig. 13-7). 94 For example, in the



Lipoprotein and Coronary Atherosclerosis Study, 13 patients with reduced HDL-C levels (on placebo) evidenced the highest rates of arteriographic progression. However, statin treatment resulted in greater decrease in progression in low versus high HDL patients that was likely a consequence of reduction in atherogenic lipoproteins and inflammation rather than raising of HDL-C. 94.95

Statins raise HDL-C levels in part by reducing CETP activity and increasing apo AI synthesis. 96 Among the different statins, rosuvastatin may raise HDL-C levels at the upper end of the spectrum (15%). 86 Similarly, simvastatin raises HDL-C and apo AI levels, 97 with the most robust increases (10% to 15%) observed at the highest dose (80 mg). 98.99

Drugs That Reduce HDL-C

Several classes of drug therapies have also been found to lower HDL-C levels. In one comprehensive review, 474 trials investigated the effects of 85 antihypertensive drugs on lipids and blood pressure. Nonselective beta blockers were shown to have a negative effect on HDL-C (10% to 20%) levels. 100,101 Subjects taking benzodiazepine derivatives have been shown to have HDL-C levels 3.3 mg/dL lower than that of nonusers. 102 However, these lipid effects may be secondary to the weight gain and accompanying increases in triglycerides. 103

Among the most potent HDL-C-lowering agents are anabolic steroids. Testosterone increases hepatic expression of hepatic lipase and SR-B1, with resulting reductions of HDL-C up to 90%. 104,105 Fortunately, this effect can be reversed within 1 month of steroid discontinuation. Taken together, beta blockers and anabolic steroids are the two classes of drugs most associated with reductions in HDL-C.

Novel Targets of High-Density Lipoprotein Metabolism

CETP Inhibition

As described before, CETP mediates the transfer of cholesteryl esters from HDL to apo B-containing lipoproteins, 14 and hereditary CETP deficiency is associated with intrinsically high total and HDL 2 cholesterol levels. Although vaccine based strategies are in development, early testing has yielded only modest increases in HDL-C (5% to 10%). 106 In contrast, several oral compounds have demonstrated more appreciable increases (> 25%). The most well studied, torcetrapib, irreversibly bound to CETP and resulted in 50% to 100% increases in HDL-C levels.

Unfortunately, the clinical trial Investigation of Lipid Level Management to Understand Its Impact in Atherosclerotic Events (ILLUMINATE) was prematurely terminated because of offtarget toxicity related to aldosterone stimulation. The increase in electrolyte abnormalities (eg, increased plasma bicarbonate and reduced potassium) was believed to contribute in part to the higher mortality rate in the torcetrapib-treated patients. 108

Another CETP inhibitor, dalcetrapib (JTT-705), regressed aortic atherosclerosis in rabbits. 109 Dalcetrapib raises HDL-C more moderately (25% to 40%) than torcetrapib does, but it does not irreversibly bind to CETP or upregulate aldosterone secretion 110 and is currently under evaluation in a large clinical outcomes trial. 111 A third compound, anacetrapib, also exhibits potent HDL-C-raising properties without affecting aldosterone. ¹¹² Pending the outcome of ongoing clinical studies, CETP inhibitors hold potential promise as adjuvant therapy in highrisk patients with dyslipidemia.



Apolipoprotein Al-Targeted Therapies

On the basis of experimental evidence that intravenous HDL infusions or apo AI overexpression reduced atherosclerosis in animals, ^{113,114} there has been great interest in human-based therapy targeting HDL. Several different strategies include administration of intravenous apo AI, reconstituted phospholipid-apo AI complexes (rHDL), oral apo AI mimetic compounds, and phospholipid-based therapy.

The first human study to gain extensive media attention was the reduction in atheroma volume (assessed by intrave nous ultrasound) after five weekly infusions of apo AI Milano/phospholipid complex in post-ACS patients. ¹¹⁵ In the randomized study Effect of rHDL on Atherosclerosis Safety and Efficacy (ERASE), ¹¹⁶ 183 ACS patients received four weekly infusions of saline or 40 mg/kg or 80 mg/kg of HDL mimetic. Although atheroma burden was not different among the groups, there was improvement in the plaque characterization index and coronary score on quantitative coronary angiography in favor of rHDL.

Apo AI mimetic peptides are much smaller compounds than mature apo AI (eg, 18 versus 243 amino acids) and possess similar apo AI/lipid-binding domains to promote cholesterol efflux, to decrease inflammation, and to improve endothelial function. $^{117}\,\mathrm{For}$ example, the mimetic D-4F improved the anti-inflammatory capacity of HDL after a single dose without altering HDL-C levels. 118

Phospholipid administration may also be a valuable HDL-targeted therapy. For example, the synthetic compound 1,2-dimyristoyl- sn-glycero-3-phosphocholine (DMPC) raised HDL and apo AI levels in association with reduced aortic lesions in a murine study. ¹¹⁹ Moreover, the soy-derived phospholipid phosphatidylinositol was found to raise HDL-C levels 13% to 18% during a 2-week period in normolipidemic subjects. ¹²⁰

Liver X Receptor Agonists

The liver X receptor (LXR), a nuclear hormone receptor (Fig. 13-9), forms a heterodimer with the retinoid X receptor and regulates transcription of ABCA1 and ABCG1, thereby serving an integral role in cholesterol efflux and reverse cholesterol transport. ¹²¹ Not surprisingly, therefore, synthetic activators of LXR have been pursued as a potential target to improve HDL functionality. Unfortunately, complicating the early-stage testing of LXR agonists was the induction of sterol

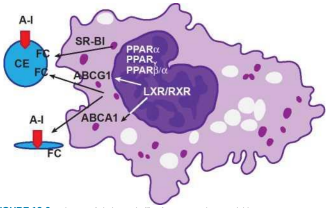


FIGURE 13-9 Pathways of cholesterol efflux from macrophages. Lipid-poor apo Al can acquire free cholesterol (FC) from macrophages through an efflux process mediated by ABCA1. Alternatively, mature HDL can promote macrophage cholesterol efflux through the ABCG1 transporter or SR-B1. ABCAI and ABCG1 expression is controlled by the nuclear receptor heterodimer LXR/RXR (liver X receptor/retinoid X receptor). The peroxisome proliferator-activated receptors (PPARs) may also influence the cholesterol efflux pathway. CE, cholesteryl ester. (From Duffy D, Rader DJ: Emerging therapies targeting high-density lipoprotein metabolism and reverse cholesterol transport. Circulation 113:1140, 2006.)

regulatory element-binding protein 1 (SREBP-1) expression, resulting in enhanced hepatic VLDL production and hypertriglyceridemia. 122

In contrast, selective targeting of macrophage LXR (ie, LXR \$\sisoform\$) was associated with reduced atherosclerosis in mice. ¹²³ Recently, a dual synthetic agonist (LXR a+P) was tested in humans and found to increase expression of ABCA1 and ABCG1 in a dose-dependent manner. However, higher doses of the compound were associated with neurological and psychiatric side effects, including confusion, forgetfulness, decreased concentration, and paranoid ideation. ¹²⁴ Although LXR-623 was withdrawn, other synthetic LXR compounds remain in clinical development.

Peroxisome Proliferator-Activated Receptor Agonists

In addition to LXR, peroxisome proliferator-activated receptors have nuclear transcription factors that also influence lipid (and glucose) homeostasis. As PPAR a agonists, fibrates upregulate apo AI transcription and promote macrophage cholesterol efflux in addition to suppressing apo C-III, resulting in increased lipoprotein lipase activity and reduced triglycerides. ⁸⁷ Other more potent PPAR a agonists are under investigation. They include the compound GFT505, which reduced triglycerides and cholesterol by 50% and inhibited aortic plaque formation, ¹²⁵ and CP-778875, which raised HDL-C up to 14% in a diabetic cohort.

The PPAR y agonists, particularly in the thiazolidinedione class, represent insulin-sensitizing agents that may raise HDL-C 5% to 15%. Of the two agents in this class, pioglitazone exerts a more favorable profile on lipids and lipoproteins, including a 15% increase in HDL-C levels compared with an 8% increase with rosiglitazone. 127 In the PROactive (PROspective Pioglitazone Clinical Trial in Macrovascular Events) study, a statistically significant difference in the primary clinical endpoint was not demonstrated. 128 However, post hoc analysis found a significant reduction in recurrent myocardial infarction and ACS in high-risk diabetic patients. 129 In contrast, the meta-analysis that suggested an increase in CHD events with rosiglitazone 130 was not confirmed in a recent multicenter trial. 131 The testing of dual combination peroxisome proliferator-activated receptor agonists, specifically of the PPAR a /PPAR y class, has been associated with adverse CHD events, leading to the discontinuation of several of these agents. In contrast, the PPAR S agonist GW501516 may hold promise in the treatment of low HDL-C and associated metabolic abnormalities. 132

Other Potential High-Density Lipoprotein Therapeutic Targets and Strategies

Two additional targets to raise HDL-C include modulation of endothelial lipase and the farnesoid X receptor (FXR). Endothelial lipase is produced by vascular endothelial cells and hydrolyzes HDL phospholipids; overexpression of endothelial lipase is associated with reduced HDL-C levels. ¹³³ Conversely , endothelial lipase inhibition is associated with increased HDL-C, ¹³⁴ as are genetic loss-of-function variants, ¹³⁵ raising the possibility that directed inhibition of endothelial lipase may be a potential HDL target.

The nuclear hormone receptor FXR is a transcriptional activator of several genes involved in the regulation of lipid metabolism. Activation of FXR results in decreased expression of SREBP-1c, a primary regulator of triglyceride biosynthesis, thereby leading to reduced triglyceride levels. However, HDL-C levels were also reduced. ¹³⁶ A natural FXR antagonist, the plant sterol guggulsterone, decreased hepatic cholesterol levels in cholesterol-fed animals, ¹³⁷ although the gugulipid extract had modest LDL-C-lowering effects without affecting HDL-C or triglycerides. ¹³⁸



213 CONCLUSION

independent inverse relationship between HDL-C levels and risk of CHD. In addition to reverse cholesterol transport, HDL possesses other atheroprotective properties, reflecting its opposition to inflammation, oxidation, and thrombosis. However, 33. Edmondson AC, Brown RJ, Kathiresan S, et al: Loss-of-function variants in endothelial lipase are the relationship between HDL-C levels and functionality is elusive, and it remains to be established whether and to what extent raising HDL-C or improving its functionality reduces CHD event rates 35. deGoma EM, deGoma RL, Rader DJ: Beyond high-density lipoprotein cholesterol levels: beyond non-HDL-based therapies. Ongoing clinical trials will, it is hoped, resolve these issues and justify whether targeting of HDL offers therapeutic value in secondary CHD prevention.

Acknowledgement

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CHAPTER 14

Low-Density Lipoprotein Cholesterol: Role in Atherosclerosis and Approaches to Therapeutic Management

Michael H. Davidson and Peter P. Toth

- Whereas LDL-C reduction has become the cornerstone of lipid management, apo B and non -HDL-C may be more accurate indicators of cardiovascular risk as well as measures by which to gauge LDL-lowering therapy.
- Clinical trials using statins and non-statin approaches to lipid lowering have demonstrated that for every 1 mg/dL reduction in serum LDL-C, there is a 1% reduction in risk for acute cardiovascular events.

KEY POINTS

- Among patients with dyslipidemia, LDL-C is the primary target of therapy. LDL-C lowering reduces risk for CHD and its clinical sequelae, including unstable angina, ischemic stroke, myocardial infarction, and death.
- Atherogenesis is a complex disease whose etiology and progression are strongly influenced by the severity of a large number of risk factors. Low serum levels of HDL-C, hypertension, impairments in glucose metabolism, heightened systemic inflammatory tone, cigarette smoking, and others induce progressive arterial wall injury.
- The metabolic syndrome is characterized by a set of five risk factors: abdominal obesity; elevated blood pressure, triglycerides, and fasting glucose concentration; and low HDL-C. Metabolic syndrome develops secondary to the effects of insulin resistance and obesity.
- A number of preventive strategies can be used to target reductions in LDL-C to decrease the burden of CHD. The NCEP ATP III guidelines identify therapeutic lifestyle changes as the initial intervention to lower LDL-C.
- Statins, or 3-hydroxy-3-methylglutaryl-coenzyme A reductase inhibitors, are the most widely used drugs for lowering of LDL-C. The statins reduce cardiovascular morbidity and mortality.
- Bile acid sequestrants, nicotinic acid, fibrates, cholesterol absorption inhibition, and omega-3 fish oils also play important roles in LDL-C reduction and dyslipidemia management.

Cardiovascular disease (CVD), including coronary heart disease (CHD), cerebrovas cular disease, hypertension, and ischemic heart disease, is a major cause of mortality and morbidity worldwide, accounting for 17 million deaths. 1 Aggressive risk reduction therapies are indicated for patients with and without CHD to improve survival, to reduce initial or recurrent events, and to improve quality of life for these patients. Among patients with dyslipidemia, low-density lipoprotein cholesterol (LDL-C) is the primary target of therapy as research has substantiated that elevated LDL-C is a major cause of morbidity and mortality and LDL-C lowering reduces the risk for CHD and its clinical including unstable sequelae, angina, myocardial infarction, and death. This chapter reviews the role of LDL-C in the development of atherosclerosis and outlines management strategies to reduce serum levels of this lipoprotein.

HISTORICAL PERSPECTIVES ON HYPERCHOLESTEROLEMIA

The importance of hypercholesterolemia in the development of atherosclerosis can be traced historically as a medical controversy, as atherosclerosis was considered an inevitable part of aging. ² Studies on rabbits demonstrating the impact of high-cholesterol diets on atherogenesis, published in 1913 by Anitschkow and Chalatov, ³ substantiated the role of hypercholesterolemia, yet it took a decade of research to confirm that the findings from experimental atherosclerosis in animals could be extrapolated to humans. ²

The Framingham Heart Study and other prospective epidemiological investigations

demonstrated that risk for myocardial infarction, stroke, and death was proportional to serum cholesterol levels. 4-6 In the Seven Countries Study, a direct relationship was found between serum cholesterol and risk for CHD among 12,763 middle -aged men. 6 This relationship has been confirmed and extended in cohorts from 26 nations in the Multinational Monitoring of Trends and Determinants in Cardiovascular Disease (MONICA) study. 7 The Ni-Hon-San study evaluated the impact of migration and progressive westernization of the diet among Japanese men residing in Nippon, Japan, or in Honolulu and San Francisco. 8 As the intake of animal fat increased, serum cholesterol and risk for CHD mortality both increased. In the Multiple Risk Factor Intervention Trial (a cohort of 356,222 men), there was a clear gradient of continuously rising risk for CVD as serum cholesterol levels increased. 9 Whereas the majority of total cholesterol is composed of LDL-C in most patients, it was important to evaluate whether LDL-C is an independent predictor of CHD. The Cooperative Lipoprotein Phe notyping study, 10 the Atherosclerosis Risk in Communities study, 11 and the Framing ham study 12 all demonstrated a direct and independent relationship between LDL-C and risk for cardiovascular morbidity and mortality. These relationships between total cholesterol and LDL-C and risk for CVD are remarkably consistent in populations studied throughout

Acceptance of the lipid hypothesis that dyslipidemia is a causative factor in the development of atherosclerosis and CHD did not begin until the early 1960s, when research findings as well as the release of the American Heart Association's recommendation for dietary modification as a first

LIPOPROTEIN METABOLISM

Cholesterol derived from both biliary and dietary sources is transported into jejunal enterocytes from the intestinal lumen by a sterol translocase known as Niemann-Pick C1-like 1 protein. ¹⁴ Cholesterol, triglycerides, and phospholipids are packaged together with apoprotein B48 to form chylomi crons. Chylomicrons enter the enteric lymphatic system and are secreted into the central circulation through the lymph duct. The triglycerides in chylomicrons can be hydrolyzed by lipoprotein lipase in serum, yielding chylomicron remnant particles. Both chylomicrons and chylomicron remnants are taken up by hepatocytes. Intrahepatic cholesterol and lipids can be repackaged with apoprotein B100 into very-low-density lipoproteins (VLDL) and secreted into the central circulation.

The triglycerides in VLDL particles are hydrolyzed by lipoprotein lipase to yield, in sequence, intermediate-density lipoproteins (IDL) and then low-density lipoproteins (LDL). LDL particles tend to be highly enriched with cholesterol. All of the apo B100-containing lipoproteins are atherogenic. The sum of all atherogenic lipoproteins (VLDL + IDL + LDL) is defined as non-high-density lipoprotein (non-HDL). Non-HDL is a fairly sensitive measure of the total atherogenic lipoprotein burden in serum. LDL particles can be taken up into arterial walls. Alternatively, LDL particles are cleared from serum by the LDL receptor (LDLR) and LDLR-related protein expressed on the surface of hepatocytes. The cholesterol in LDL particles can then be shunted towards bile acid formation (rate-limiting step catalyzed by 7 a -hydroxylase), it can be secreted into bile, or it can be repackaged into VLDL.

In patients with loss-of-function mutations in lipoprotein lipase, serum VLDL and triglyceride levels tend to be elevated. This impairs the conversion of VLDL into smaller, triglyceride-depleted lipoproteins. With this excess availability of triglyceride in VLDL, cholesteryl ester transfer protein catalyzes a 1:1 stoichiometric exchange for cholesterol between VLDL and LDL. This leads to progressive enrichment of the LDL particles with triglyceride, rendering them more vulnerable to lipolysis by the hepatic lipase enzyme. Hepatic lipase converts large buoyant LDL particles into smaller, denser and more numerous particles. Smaller LDL particles also tend to be more atherogenic: (1) biophysically, they can gain access into the subendothelial space more easily because they are smaller; (2) they are more easily oxidized than their larger counterparts; and (3) they have a lower affinity for the LDLR and hence reduced systemic clearance.

Some patients have a genetic predisposition to elevated serum levels of LDL-C. Hereditary hypercholesterolemias are frequently associated with premature multivessel coronary artery disease (CAD). Patients with familial hypercholesterolemia harbor a variety of mutations (these are indexed at www.ucl.ac.uk/fh/) in the gene for LDLR, resulting in reduced capacity for LDL clearance from serum. 15 Heterozygotes often present with serum LDL-C of 250 to 350 mg/dL. Homozygotes can have LDL-C levels of 500 mg/dL or more and usually require LDL apheresis to control their serum levels of this lipoprotein.

Loss-of-function mutations in cholesterol 7 a -hydroxylase are also associated with elevations in LDL-C and decreased bile acid

biosynthesis. ¹⁶ Patients with familial defective apo B100 have impaired capacity for LDL-C clearance secondary to polymorphisms in apo B100 (henceforth apo B), which reduce the affinity of this apoprotein for the LDLR. ¹⁷ Another fascinating group of mutations linked to hypercholesterolemia localizes to the gene coding for the enzyme proprotein convertase subtilisin/kexin type 9 (PCSK9). ^{18,19} This enzyme modulates the activity of the LDLR. Gain-of-function mutations lead to increased degradation of the LDLR, whereas loss-of-function mutations allow increased expression of the LDLR; these mutations are associated with increased and decreased risk for CAD, respectively.

LOW-DENSITY LIPOPROTEIN CHOLESTEROL AND ATHEROSCLEROTIC VASCULAR DISEASES

Atherogenesis is a complex disease whose etiology and progression are strongly influenced by the severity of a large number of risk factors. LDL is an apo B-containing particle (Fig. 14-1) and as such contributes to high levels of atherogenic lipoproteins (Fig. 14-2). In addition, low serum levels of highdensity lipoprotein cholesterol (HDL-C), hypertension, impairments in glucose metabolism, heightened systemic inflammatory tone, cigarette smoking, and other risk factors induce progressive arterial wall injury. An early event in atherogenesis is the development of endothelial cell dysfunction . Under normal physiological conditions, the endothelium serves as an efficient barrier between blood and the arterial wall. The endothelium produces nitric oxide (a potent vasodilator) and maintains an antithrombogenic surface by producing prostacyclin and tissue plasminogen activator. As the endothelium becomes stressed in response to risk factors, it becomes dysfunctional, resulting in a maladaptive inflammatory response (Fig. 14-3). 20,21

Endothelial dysfunction is highly associated with atherosclerosis. Dysfunctional endothelial cells increase the expression of adhesion molecules, such as vascular cell adhesion molecule 1, intercellular adhesion molecule 1, and different types of selectins. ²² These cell surface receptors interact with



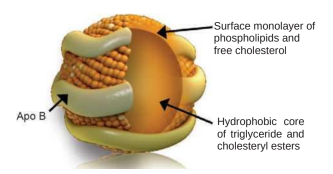
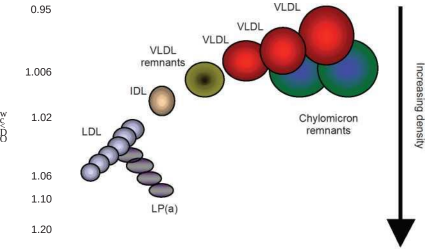


FIGURE 14-1 The structure of low-density lipoprotein. Nuclear magnetic resonance (NMR) spectroscopy of the LDL surface structure has enabled identification of a surface monolayer of phospholipids and free cholesterol and a hydrophobic core of triglyceride and cholesteryl esters. Phosphatidylcholine and sphingomyelin are tightly bound to apo B and therefore NMR invisible. (From Murphy HC, Burns SP, White JJ, et al: Investigation of human low density lipoprotein by 1H nuclear magnetic resonance spectroscopy: mobility of phosphatidylcholine and sphingomyelin headgroups characterizes the surface layer. Biochemistry 39:9763, 2000. Reproduced with permission.)

FIGURE 14-2 The atherogenicity of apo B-containing lipoproteins is related to their diameter and density. Chylomicrons are the largest lipoproteins and are produced by jejunal enterocytes. VLDL is secreted by the liver and progressively converted into IDL and then LDL by the activity of serum lipases. Atherogenic lipoproteins include chylomicron remnants, VLDL, VLDL remnants, IDL, and LDL particles. Remnants represent particles that are incompletely hydrolyzed. Lipoprotein(a) [Lp(a)] is a variant of LDL that is also atherogenic. Dyslipoproteinemia with low concentrations of HDL-C and elevated serum triglycerides with increased concentrations of small, dense LDL particles is associated with a particularly high incidence of coronary vascular disease. (From Segrest JP, Garber DW, Brouillette CG, et al: The amphipathic alpha helix: a multifunctional structural motif in plasma apolipoproteins. Adv Protein Chem 45:303, 1994. Reproduced with permission.)



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counterreceptors on the surface of inflammatory white blood cells, such as monocytes, T cells, and mast cells. These receptor-counterreceptor interactions facilitate the binding, rolling, and stable arrest of these cells along the endothelial surface.

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Under normal conditions, endothelial cells maintain appropriate intercellular contacts through gap junctions, which allow these cells to communicate with each other and to maintain tight contacts among themselves. ²³ These gap junctions also become dysfunctional and allow the transmi gration of inflammatory white cells into the subendothelial space by following a gradient of monocyte chemoattractant protein 1 down into the subendothelial space. Once they are in the subendothelial space, monocytes can transform into resident macrophages in response to macrophage colony stimulating factor. As gap junctions and endothelial cells become more dysfunctional, their barrier function becomes progressively more compromised, allowing greater influx of atherogenic lipoproteins into the arterial wall. Lipoproteins can be trapped in the vessel wall by intercellular matrix proteins such as proteoglycans, fibronectin, and vitronectin, among others. 24

In the subendothelial space, LDL particles can be oxidized by such enzymes as myeloperoxidase, 5' - lipoxygenase, lipo protein-associated phospholipase A 2, NADPH oxidase, and others. ^{25,26} T cells can present oxidized LDL to macrophages. The macrophages are activated and increase the expression of families of scavenging receptors (eg, CD36, scavenging receptor A). Scavenging receptors take up oxidized LDL, leading to foam cell formation. Foam cells can coalesce to form fatty streaks, and fatty streaks are the progenitors to the formation of atherosclerotic plaque. Macrophages can maintain a certain level of lipid homeostasis as long as there is adequate availability of HDL particles. HDL particles interact with macrophages and foam cells to promote the mobilization and externalization of intracellular lipid. ²⁷⁻²⁹

If there is inadequate availability of HDL in the face of excess LDL in the extracellular milieu, there is continued net accumulation of lipid by the macrophage, ultimately resulting in apoptosis or programmed cell death. As cellular debris and more lipid accumulate, and as the capacity to phagocytose and clear this debris is progressively more compromised, the volume of the atherosclerotic plaque increases and develops a lipid core ³⁰ (Fig. 14-4).

Diameter (nm)

20 40 60 80 NATIONAL CHOLESTEROL EDUCATION PROGRAM GUIDELINES

A high serum level of LDL-C is an established risk factor for the development of CHD. For this reason, LDL-C reduction is the primary goal of therapy in patients with dyslipidemia. This recommendation is emphasized by the National Choles terol Education Program Adult Treatment Panel III (NCEP ATP III), 31 American Heart Association/American College of Cardiology guidelines for secondary prevention of CHD, 32 and the European Guidelines on Cardiovascular Disease Prevention. 33 NCEP ATP III defines an LDL-C of < 100 mg/dL as optimal. Among patients in the primary prevention setting, it is recommended that the 10-year projected risk for sustaining a CHD-related event be estimated with the Framingham risk score. Non-HDL-C is a surrogate measure of total atherogenic lipoprotein burden in serum and represents the sum of VLDL, IDL, LDL, and lipoprotein(a) as well as remnant particles. It is calculated by subtracting HDL-C from total cholesterol. Among patients with baseline triglycerides > 200 mg/dL, non-HDL-C is the secondary target of therapy. Non-HDL-C has been shown to be a better predictor than LDL-C of risk for cardiovascular events in both men and women. 34,35 Goals for both LDL-C and non-HDL-C are risk stratified, and the goals for both of these lipoprotein fractions decrease as risk increases (Table 14-1). Among patients with CAD or a CAD risk equivalent (diabetes mellitus, abdominal aortic aneurysm, peripheral arterial disease, symptomatic carotid artery disease, or 10-year Framingham risk score > 20%), it is recommended that patients achieve LDL-C < 100 mg/dL and non-HDL-C < 130 mg/dL. Among patients defined as very high risk (CAD complicated by a recent acute coronary syndrome, diabetes mellitus, or multiple poorly controlled risk factors), it is a therapeutic option to reduce LDL-C below 70 mg/dL and non-HDL-C below 100 mg/dL. 31 In a more recent advisory statement from the American Heart Association, it was recommended that an LDL-C target of < 70 mg/dL be an option for any patient with established CAD. 36

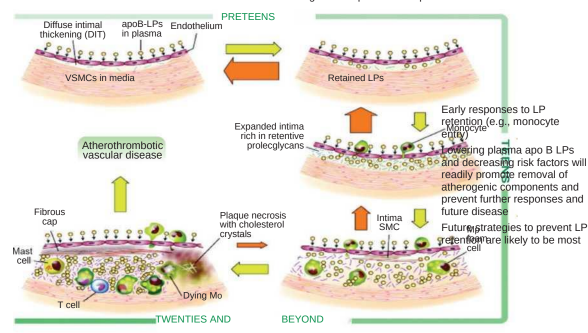
Evidence from clinical trials has established the merits of aggressive LDL-C risk reduction therapies for patients with CHD and other atherosclerotic vascular diseases in improving survival, reducing recurrent coronary events and need for revascularization, and improving quality of life. Date from

Prelesional susceptible area of the arterial wall with diffuse intimal thickening (DIT)

Lowering plasma apo B LPs and decreasing risk factors will prevent future vascular disease

vascular disease may still develop Early lipoprotein retention

Lowering plasma apo B LPs and decreasing risk factors will readily promote removal of atherogenic components and prevent



- Advanced responses to LP retention, including maladaptive inflammation, MP death, and plaque necrosis
- · LP retention continues to accelerate
- Lowering plasma apo B LPs and reducing risk factors can promote removal of atherogenic components and promote regression, but reversal is more difficult and prolonged, and

maladaptive responses and future disease

- Continued responses to LP retention, (eg, MP foam cell formation and SMC migration)
- · LP retention starts to accelerate
- Lowering plasma apo B LPs and other risk factors can still promote removal of atherogenic components, promote regression, and prevent further responses and future disease

FIGURE 14-3 The continuum of developing atherosclerotic disease over a lifetime. Atherogenesis involves a complex interplay between inflammation, apo B-containing lipoproteins, and histological components of blood vessel walls. Atherogenesis is potentiated by apo B lipoprotein retention by subendothelial extracellular matrix molecules (eg. proteoglycans), which initiates a series of biological reactions that develop into a maladaptive inflammatory response. As depicted in this figure, early response to lipoprotein retention begins in the preteen years and continues to accelerate in the 20s and beyond. Green arrows indicate progression; orange arrows indicate the potential for regression. The earliest stages are the most easily reversible by lowering of plasma apo B lipoproteins (large orange arrows). The complexity of advanced lesions, including accelerated lipoprotein retention, renders them less reversible (small orange arrows) as the severity of atherosclerosis advances. (From Tabas I, Williams KJ, Boren J: Subendothelial lipoprotein retention as the initiating process in atherosclerosis: update and therapeutic implications. Circulation 116:1832, 2007. Reproduced with permission.)

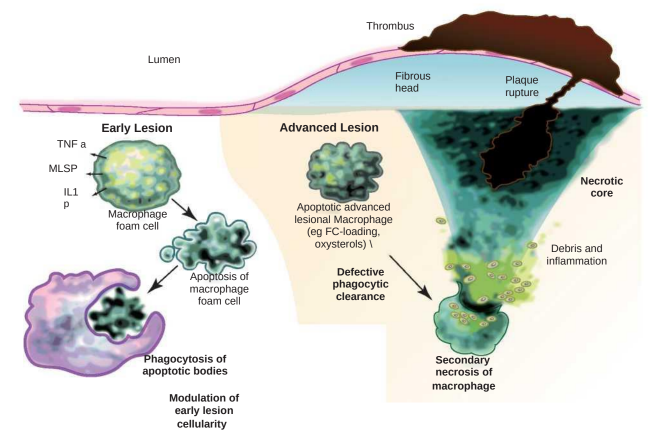


FIGURE 14-4 The role of macrophage apoptosis and atherosclerotic lesion progression. In the setting of endothelial dysfunction, atherogenic lipoproteins enter the subendothelial space, where they can aggregate and become oxidatively modified by a variety of enzymes. Oxidized LDL induces the upregulation of scavenging receptors on the surface of macrophages resident in the subendothelial space. Progressive macrophage lipid uptake results in the formation of foam cells. There is a limit to lipid uptake. Excess lipid is toxic to the cell and can result in apoptosis. Early in the course of atherogenesis, apoptotic debris is phagocytosed by other macrophages, a process that can modulate the cellularity and rate of progression of an atherosclerotic lesion. As depicted in this figure, in late lesions (right), macrophages also undergo apoptosis but are no longer cleared as efficiently. This leads to the formation of the necrotic core, which promotes inflammation, plaque instability, and acute lesional thrombosis secondary to loss of architectural integrity and rupture. (From Tabas I: Consequences and therapeutic implications of macrophage apoptosis in atherosclerosis: the importance of lesion stage and phagocytic efficiency. Arterioscler Thromb Vasc Biol 25:2255, 2005. Reproduced with permission.)

TABLE 14—1 LDL-C Goals and Three	esholds for Initiation of Lifestyle	Change and Pharmacologic Intervention	
Risk Category ¹	LDL Goal	LDL Level at Which to Initiate TLC	LDL Level at Which to Consider Drug Therapy
CHD or CHD risk equivalents (10-year risk > 20%)	< 100 mg/dL (optional blank < 70) ²	> 100 mg/dL All patients regardless of LDL	> 130 mg/dL (100-129 mg/dL: drug of choice) > 100 mg/dL · (< 100 mg/dL: drug of choice)
2 + risk factors (10-year risk 10%-20%)	< 130 mg/dL (optional blank < 100)	> 130 mg/dL All patients regardless of LDL	> 130 mg/dL (> 100 mg/dL: optional drug ⁽)
2 + risk factors (10-year risk < 10%)	< 130 mg/dL	> 130 mg/dL	> 160 mg/dL
0-1 risk factor	< 160 mg/dL	> 160 mg/dL	> 190 mg/dL (160-189 mg/dL: LDL-lowering drug optional)

Coronary heart disease (CHD) risk equivalents include diabetes mellitus, peripheral vascular disease, carotid artery disease, and abdominal aortic aneurysm. TLC, therapeutic lifestyle changes.

¹ Risk factors included in Framingham risk evaluation are age, systolic blood pressure, total cholesterol, HDL-C, and smoking status.

²The optional goal of <70 mg/dL is particularly targeted at patients who have "very high" risk (eg, patients with a recent acute coronary syndrome, poorly controlled diabetics with multiple risk factors).

³When statin therapy is initiated in these patients, the goal for LDL-C reduction should be 30% to 40% from baseline.

Based on Grundy SM, Cleeman JI, Merz CN, et al: Implications of recent clinical trials for the National Cholesterol Education Program Adult Treatment Panel III guidelines. Circulation 110:227, 2004.

However, despite the substantial body of research establishing the benefit of LDL-C reduction, only a small percentage of patients with CHD are reaching lipid goals. In the National Cholesterol Education Program (NCEP) Evaluation Project Utilizing Novel E-Technology (NEPTUNE) survey of physicians regarding the care of 4885 patients with dyslipidemia ,75% of patients with CHD met the definition of "very high risk," yet only 18% had an LDL-C level < 70 mg/dL and only 4% had an LDL-C level < 70 mg/dL and a non-HDL-C level < 100 mg/dL when triglycerides were > 200 mg/dL. $^{\rm 42}$

In a retrospective study from a national outpatient electronic medical record database of 10,637 patients with a diagnosis of atherosclerosis, 57% of the 4067 patients with a baseline LDL-C > 100 mg/dL were not prescribed statin treatment after diagnosis. Among patients receiving statin or any other cholesterollowering therapy after diagnosis who had baseline and followup LDL-C values (n = 682), 43% had a post-diagnosis LDL-C > 100 mg/dL. ⁴³ The proportion of patients with LDL-C < 100 mg/dL at baseline was 46.8%, and 12% of patients had LDL-C cholesterol < 70 mg/dL. Less than 5% of patients were currently receiving hypercholesterolemia therapy at the time of diagnosis, and 25% had hypercholesterolemia treatment after their condition had been diagnosed. Among patients receiving hypercholesterolemia therapy, LDL-C levels in 12 months after diagnosis of atherosclerosis were similar to levels at the time of diagnosis. These data substantiate the need for more aggressive statin therapy and implementation of combination therapy to reduce residual risk for cardiovascular events and highlight the importance of monitoring and managing lipid levels in patients with atherosclerosis.

METABOLIC SYNDROME AND CARDIOMETABOLIC RISK

In the United States, it is estimated that 35% to 40% of adults have the metabolic syndrome, a cluster of lipid and nonlipid disorders. 44,45 The metabolic syndrome is characterized by a set of five risk factors: abdominal obesity; elevated blood pressure, triglycerides, and fasting glucose concentration; and low HDL-C (Table 14-2). The presence of any three or more of the following risk factors in a single individual is diagnostic of the metabolic syndrome according to the revised ATP III/American Heart Association/National Heart, Lung, and Blood Institute definition 46 : triglycerides > 150 mg/ dL, HDL-C < 40 mg/dL in men and < 50 mg/dL in women, serum glucose concentration > 100 mg/dL, blood pressure > 130/85 mm Hg (can be either systolic or diastolic or on therapy with antihypertensive medication), and waist circumference > 40 inches in men and > 35 inches in women. Metabolic syndrome develops secondary to the effects of insulin resistance and obesity. Although the metabolic syndrome significantly increases the risk for atherosclerotic disease and diabetes mellitus, it is not defined as a CAD risk equivalent.

The primary goal of clinical management of the metabolic syndrome is to reduce CHD risk and diabetes mellitus. ³¹ This focuses on intensified LDL-C lowering and modification of the underlying risk factors including obesity, physical inactivity , and other risk factors associated with the metabolic

Risk Factor	Defining Level
Abdominal obesity Men Women Triglycerides	Waist > 40 inches Waist > 35 inches > 150 mg/dL
HDL-C	
Men Women	< 40 mg/dL < 50 mg/dL
Blood pressure	> 130/ > 85 mm Hg or taking antihypertensive medication

*Patients having any three of the five risk factors meet the criteria for the diagnosis of the metabolic syndrome.

> 100 mg/dL or taking hypoglycemic medication

Modified from Toth P: Dyslipoproteinemias. *In* Rakel RE, Bope ET, editors: *Cohn's current therapy,* Philadelphia, 2006, Elsevier.

Fasting glucose

TABLE 14-3 Dietary Recommendations for Therapeutic Lifestyle Change				
Dietary Component	Recommendation Allowance			
Polyunsaturated fat	Up to 10% of total calories			
Monounsaturated fat	Up to 20% of total calories			
Totally done	25%-35% of total calories			
Carbohydrates	50%-60% of total calories			
Dietary fiber	20-30 g/day			
Protein	Approximately 15% of total calories			
Dietary cholesterol	< 200 mg/day			

syndrome, such as blood pressure control (Table 14-3). 31 The NCEP ATP III guidelines emphasize the modification of metabolic syndrome risk factors through lifestyle changes, especially because both weight loss and exercise reduce insulin resistance and favorably modify risk factors for the metabolic syndrome. 31

THERAPEUTIC LIFESTYLE CHANGES

A number of preventive strategies can be used to target reductions in LDL-C to decrease the burden of CHD. The NCEP ATP III guidelines identify therapeutic lifestyle changes (TLC) as the initial intervention for lowering LDL-C with a focus on smoking cessation, weight loss, total calories, physical activity to maintain desirable weight, and moderate alcohol intake (see Table 14-3). The NCEP guidelines outline TLC as a multifaceted lifestyle approach for LDL-C lowering. The recommendations for TLC include a reduction in satu rated fat intake to < 7% of total calories and cholesterol to < 200 mg/day, weight reduction, increased physical activity, therapeutic options to enhance LDL lowering such as the use of plant stanols or sterols (2 g/day), and increased viscous (soluble) fiber (10 to 25 g/day) to reduce cholesterol absorption. ³¹ The aim of primary prevention is to reduce long-term risk (> 10 years) as well as short-term risk (< 10 years). Whereas TLC is the basis of clinical primary prevention, pharmaco logical therapy is often indicated in the management of ele vated LDL-C to achieve risk-stratified target



STATIN THERAPY

Statins, or 3-hydroxy-3-methylglutaryl-coenzyme A reduc tase inhibitors, have been the most widely used therapy for the treatment of dyslipidemia to reduce the risk of CHD. Statins are used to target the reduction of elevated LDL-C and to improve the lipid level profile. The statins are recognized as the first-line treatment of dyslipidemia. 31-33,47 Statins are used to target the reduction of elevated LDL-C and to improve all components of the lipid level profile.

Data from the major statin trials including the Scandinavian Simvastatin Survival Study (4S), ⁴⁸ Cholesterol and Recurrent Events (CARE), ⁴⁹ HPS, ³⁹ Long-term Intervention with Pravastatin in Ischemic Disease (LIPID), ³⁸ and TNT ⁴⁰ have established that LDL-C reduction correlates with cardio vascular event reduction in a linear fashion. The HPS, the largest statin trial, demonstrated the benefits of aggressive

statin therapy with simvastatin 40 mg/day, with a 24% **221** reduction in CVD events compared with placebo (Table 14-4). The beneficial effects of LDL-C reduction were demonstrated even in patients with baseline LDL-C levels below 100 mg/dL and in persons with diabetes.

The TNT trial additionally validated the benefit of lowering LDL-C to < 100 mg/dL with treatment with atorvastatin 80 mg daily, resulting in a 22% reduction of cardiovascular events compared with atorvastatin 20 mg daily. The IDEAL 41 study established the benefit of intensive lipid lowering with statin therapy with atorvastatin 10 mg daily, resulting in a lower LDL-C level and an 11% reduction in major cardiovascular events compared with simvastatin 20 mg daily. A meta analysis of 14 prospective randomized statin trials by the Cholesterol Treatment Trialists Collaboration showed that for every 39 mg/dL (1 mmol/L) reduction in serum LDL-C with statin therapy during a mean 5-year follow-up, there was a

		Primary Prevention Studies	
Study AFCAPS/TexCAPS	bar Lovastatin 20 to 40 mg/day versus	Design 6605 men and women	Outcomes
	placebo		40% reduction in fatal and non-fatal MI; 37% reduction in firs ACS; 33% reduction in coronary revascularizations; and unstable angina reduced by 32%
ASCOT	Atorvastatin 10 mg/day versus		
	placebo	10,305 hypertensive men (n = 8463) and women (n = 1942) with treated high blood pressure and no previous CAD	36% reduction in total CHD/nonfatal MI; 27% reduction in fatal and non-fatal strokes; total coronary event reduced by 29%; fatal and nonfatal strokes reduced by 27%
CARDS	Atorvastatin 10 mg/day versus placebo	2838 patients with type 2 diabetes mellitus and 1 CHD risk factor	37% reduction of major cardiovascular events; 27% reduction of total mortality; 13.4% reduction of acute cardiovascular events; 36% reduction of acute coronary events; 48% reduction of strokes
Heart Protection Study	Simvastatin 40 mg/day versus placebo	20,536 high-risk individuals (previous CHD, other vascular disease, hypertension among men aged > 65 years, or diabetes)	25% reduction in all-cause and coronary death rates and in strokes; need for revascularization reduced by 24%; fatal and nonfatal stroke reduced by 25%; nonfatal MI reduced by 38%; coronary mortality reduced by 18%; all-cause mortality reduced by 13%; cardiovascular event rate reduced by 24%
PROSPEROUS	Pravastatin 40 mg/day versus placebo	5804 men (n = 2804) and women (n = 3000) aged 70 to 82 years	15% reduction in combined endpoint (fatal/nonfatal MI or stroke); 19% reduction in total/non-fatal CHD; no effect on stroke (but 25% reduction in TIA)
WOSCOPS	Pravachol therapy 40 mg/day versus placebo	6595 men	CHD death of nonfatal MI reduced by 31%; CVD death reduced by 32%; total mortality 22% reduction
Study 4S	bar Simvastatin 20 mg/day versus	Secondary Prevention Studies Design 4444 patients with angina pectoris or history	Outcomes
	placebo	of MI	Coronary mortality reduced by 42%; myocardial revascularization reduction of 37%; all-cause mortality reduced by 30%; nonfatal major coronary event reduced by 34%; fatal and nonfatal stroke reduced by 30%
WARNING	Atorvastatin 80 mg/day versus angioplasty + usual care	341 patients with stable CAD	36% reduction in ischemic events; delayed time to first ischemic event reduced by 36%
WHICH	Pravastatin 40 mg/day versus placebo	3583 men and 576 women with history of MI	Death from CHD or nonfatal MI reduced by 24%; death from CHD reduced by 20%; nonfatal MI reduced by 23%; fatal MI reduced by 37%; CABG or PTCA reduced by 27%
IDEAL	Atorvastatin 80 mg/day versus simvastatin 20-40 mg/day	8888 men and women with CHD	Major cardiac events reduced by 13%; nonfatal MI reduced by 17%; revascularization reduced by 23%; peripheral arterial disease reduced by 24%

Continued



Study JUPITER	bar Rosuvastatin 20 mg/day versus	Secondary Prevention Studies Design	Outcomes
	placebo	17,802 men (> 50 years) and women (> 60 years) with no history of CAD or diabetes mellitus, entry LDL < 130 mg/dL, and CRP > 2.0 mg/L	44% reduction in primary endpoint of major coronary events; 65% reduction in non-fatal MI; 48% reduction in non-fatal strokes; 46% reduction in need for revascularization; 20% reduction in all-cause mortality
LIPID	Pravachol 40 mg/day versus placebo	9014 patients	Coronary mortality reduced by 24%; strokes reduced by 19%; fatal CHD or nonfatal MI reduced by 24%; fatal or nonfatal MI reduced by 29%
shortage	Fluvastatin 40 mg/day versus placebo	1667 men and women aged 18-80 years after angioplasty for CAD	22% lower rate of major coronary events (eg, cardiac deaths, nonfatal MI, or reintervention procedure)
MIRACLE	Atorvastatin 80 mg/day versus placebo	3086 patients with ACS	Reduction in composite endpoint by 16%; ischemia reduced by 26%; stroke reduced by 50%
PROVE IT	Atorvastatin 80 mg/day versus pravastatin 40 mg/day	4162 patients with ACS	16% reduction of composite endpoint; 14% reduction in CHD death, MI, or revascularization; revascularizations reduced by 14%; unstable angina reduced by 29%
REVERSE	Atorvastatin 80 mg/day versus pravastatin 40 mg/day	654 patients with CAD	Atheroma: atorvastatin - 0.4%, pravastatin 2.7%, difference of - 3.1%, $P = 0.02$
TNT	Atorvastatin 10 mg/day versus 80 mg/day	10,003 patients with CHD and LDL cholesterol 130-250 mg/dL	22% reduction in composite endpoint; MI reduced by 22%; stroke reduced by 25%

ACS, acute coronary syndrome; CABG, coronary artery bypass grafting; CAD, coronary artery disease; CHD, coronary heart disease; CRP, C-reactive protein; LDL, low-density lipoprotein: MI, myocardial infarction: PTCA, percutaneous transluminal coronary angioplasty: TIA, transient ischemic attack

AFCAPS/TexCAPS, Air Force/Texas Coronary Atherosclerosis Prevention Study: Implications for Preventive Cardiology in the General Adult US Population (Curr Atheroscler Rep 1:38, 1999); ASCOT, Anglo-Scandinavian Cardiac Outcomes Trial—Lipid Lowering Arm (Lancet 361:1149, 2003); CARDS, Collaborative Atorvastatin Diabetes Study (Lancet 364:685, 2004); Heart Protection Study (Lancet 360:7, 2002); PROSPER, Pravastatin in elderly individuals at risk of vascular disease (Lancet 360:1623, 2002); WOSCOPS, West of Scotland Coronary Prevention Study (N Engl J Med 333:1301, 1995); 4S, Scandinavian Simvastatin Survival Study (Lancet 344:1383, 1994); AVERT, Atorvastatin versus Revascularization Treatment Investigators (N Engl J Med 341:70, 1999); CARE, Cholesterol and Recurrent Events Trial (N Engl J Med 335:1001, 1996); IDEAL, Incremental Decrease in End Points Through Aggressive Lipid Lowering Study (JAMA 294:2437, 2005); JUPITER: The Justification for the Use of Statins in Prevention: an Intervention Trial Evaluating Rosuvastatin (N Engl J Med 359:2195, 2008); LIPID, Long-Term Intervention with Pravastatin in Ischemic Disease (Am J Cardiol 76:474, 1995); LIPS, Lescol Intervention Prevention Study (JAMA 287:3215, 2002); MIRACL, Myocardial Ischemia Reduction with Aggressive Cholesterol Lowering Study (JAMA 285:1711, 2001); PROVE IT, Pravastatin or Atorvastatin Evaluation and Infection Therapy Study (N Engl J Med 350:1495, 2004); REVERSAL, The REVERSING Atherosclerosis with Aggressive Lipid Lowering Study (JAMA 292:1, 2004); TNT, Treating to New Targets trial (N Engl J Med 352:1425, 2005).

From Toth PP: Management of dyslipidemia. In Toth PP, Cannon CP, editors: Comprehensive cardiovascular medicine in the primary care setting, Philadelphia, 2010, Humana-Springer. Reproduced with permission.

12% reduction in all-cause mortality, a 19% reduction in coronary mortality, a 24% reduction in myocardial infarction or coronary death, a 24% reduction in need for revascularization, and a 17% reduction in fatal or nonfatal stroke. ⁵⁰

The predominant benefits of statins on CHD risk reduction are due to their LDL-C-lowering effects, but the effects that statins have on triglycerides and HDL-C may also contribute to risk reduction. In the 4S 48 and The Air Force/Texas Coro nary Atherosclerosis Prevention (AFCAPS/TexCAPS) 51 trial, apo B rather than LDL-C was a better predictor of event reduction . As apo B is contained in LDL-C, the triglyceride-containing IDL, and VLDL, the impact of statins on IDL may also affect risk reduction.

The AFCAPS/TexCAPS trial additionally demonstrated that the best overall predictor of events on treatment was the ratio apo B/apo AI. This supports the concept that raising HDL (the apo AI-containing particles) also contributes to the riskmodifying benefits of statin therapy. The Justification for the Use of statins in Prevention: an Intervention Trial Evalu ating Rosuvastatin (JUPITER) 52 trial demonstrated the benefit of statin therapy on dual targets of LDL-C and high-sensitivity C-reactive protein (hsCRP), with subjects receiving rosuvas tatin 20 mg/day who achieved both LDL-C < 70 mg/dL and hsCRP < 2.0mg/L having the lowest incidence of vascular events. The JUPITER trial also provided compelling evidence that ontreatment LDL-C level is a strong predictor of clinical benefit. Compared with placebo, participants allocated to rosuvastatin who did not achieve LDL-C < 70 mg/dL had no significant reduction in vascular events (placebo event rate, 1.11%/year; participant allocated to rosuvastatin with LDL-C > 70 mg/dL event rate, 0.91%/year; HR, 0.89; 95% CI, 0.65 125; P = 0.49). Alternatively, patients allocated to rosuvastatin who had ontreatment hsCRP > 2.0 mg/L still had a significant benefit (event rate, 0.7%/year), but not as much as those who achieved an hsCRP < 2.0 mg/L (event rate, 0.42%/year).

Achievement of the dual goals of LDL-C < 70 mg/dL and CRP < 2.0 mg/L is difficult as demonstrated in a 2-year follow up of the Pravastatin Or atorVastatin Evaluation and Infection Therapy (PROVE-IT) study, which tested the efficacy of ator vastatin 80 mg/day and pravastatin 40 mg/day (Fig. 14-5). Whereas atorvastatin 80 mg was superior to pravastatin 40 mg/day in terms of achieving the dual goals of aggressive LDL-C and CRP reduction, neither agent brought the majority of patients below thresholds needed to maximize risk reduction. 53

Lipid changes appear to explain the expected outcome benefits of statin therapy. The pleiotropic or added effects of statins, such as vasodilation, antithrombosis, antioxidant, antiproliferative, and anti-inflammatory effects, may also play a role in CHD risk reduction, although this is not yet firmly established. A recent meta-analysis highlighted the

ACS population - 2 year F/U - Pravastatin 40 mg vs Atorvastatin 80 mg

myocardial infarction or

0.0

0.5 1.0 1.5 2.0

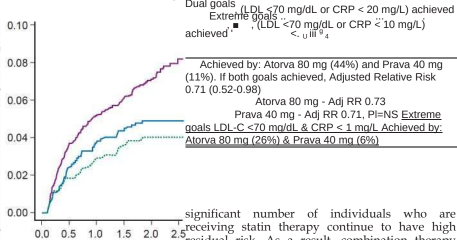
FIGURE 14-5 In a 2-year follow-up study comparing the efficacy of statin regimens in achieving the dual goals of low-density lipoprotein cholesterol (LDL-C) and C-reactive protein (CRP) reduction, 44% of patients receiving atorvastatin 80 mg/day achieved the dual goals, whereas 11% of patients receiving pravastatin 40 mg/day achieved the dual goals. As shown in this figure, both groups achieved similar risk reduction, but neither agent brought the majority of patients below thresholds needed to maximize benefit, substantiating that the dual goals of LDL-C < 70 mg/dL and CRP < 2.0 mg/L are difficult to achieve. (From Ridker PM, Morrow DA, Rose LM, et al: Relative efficacy of atorvastatin 80 mg and pravastatin 40 mg in achieving the dual goals of low-density lipoprotein cholesterol <70 mg/dl and C-

reactive protein < 2 mg/l. J Am Coll Cardiol 45:1644,

2005. Reproduced with permission.)

- Dual goals LDL-C <70 mg/dL and CRP<2.0 mg/L hard to achieve
- Risk reduction the same regardless of method Dual goals (LDI

Prove It



association of LDL-C reduction and incidence of stroke among the major statin trials, further demonstrating the beneficial effects of statin therapy in CHD risk reduction. 54

STATIN SAFETY

The safety of high-dose statin therapy is a clinical concern as associated adverse events include statin-associated persistent liver enzyme elevation, myotoxicity, and life-threatening rhabdomyolysis. Whereas the reported occurrence of musclerelated adverse events in clinical trials of statins is low, many of the trials were not designed or powered to detect rhabdo myolysis. Simultaneously, different definitions of adverse events have been used in the various clinical trials. A standard definition of nonfatal rhabdomyolysis is the development of muscle symptoms plus creatine kinase > 10 times the upper limit of normal. In terms of incidence rates of rhabdo myolysis, in the HPS, ³⁹ which was the largest clinical trial of statin therapy to date, there were five cases (0.05%) of non fatal rhabdomyolysis in patients receiving simvastatin 40 mg compared with three cases (0.03%) in patients receiving placebo. Despite the fact that the absolute event rate for rhabdomyolysis remains low at all doses for approved statins, the highest approved dose of a statin does have an increased risk for muscle adverse effects. There does not, however, appear to be a relationship between risk for myopathy and achieved level of LDL-C. 40,50 It therefore becomes necessary to acknowledge the risk of statin therapy even in standard doses used to reduce cardiovascular risk. Patients should be counseled about symptoms associated with skeletal muscle and liver toxicity.

COMBINATION THERAPY

not achieved

Whereas statin therapy is an established strategy of reducing death and myocardial infarction among patients with CHD, a

Dual goals , (LDL > 70 mg/dL or CRP > 20 mg/L)

residual risk. As a result, combination therapy Follow-up (years) increases the likelihood of achieving target lipid levels, especially in those patients with residual risk. Combination therapy results in a greater degree of reduction in residual risk of ĈVD events compared with monotherapy. Reasons for considering combination therapy include lack of achievement of LDL goals, lack of achievement of non-HDL-C goals, inadequate HDL-C elevation, and safety concerns associated with the use of high doses of statins.

Bile acid sequestrants (cholestyramine, colestipol, and colesevelam hydrochloride), niacin (extended release), and ezetimibe lower LDL-C by 15% to 20%; a 15% to 20% decrease in LDL-C is approximately equivalent to tripling the dose of a statin. For every doubling of the dose of a statin, there is a further decrease in LDL-C of approximately 6%. Therefore, the use of combination therapy is indicated to further improve the lipid profile, especially for very high risk patients who have not yet achieved the optional therapeutic target. The NCEP ATP III guidelines indicate that statin combination therapy with a bile acid sequestrant or nicotinic acid is indicated if LDL-C goals are not achieved. 31

STATINS-BILE ACID SEQUESTRANTS

Bile acid sequestrants are a class of antihyperlipidemic drugs that augment cholesterol excretion through enhanced conversion to bile acids to lower LDL-C. By reducing intrahepatic stores of cholesterol, bile acid sequestrants stimulate increased expression of the LDLR and promote increased clearance of LDL-C. In the Lipid Research Clinics Coronary Primary Prevention Trial, lipid lowering with cholestyramine was associated with a significant 19% reduction in the composite endpoint of nonfatal myocardial infarction and death among men with no prior history of CHD.

Two of the most commonly prescribed bile acid sequestrants, cholestyramine and colestipol, have established

efficacy and safety as nonsystemic approaches to cholesterol reduction. Several large-scale clinical trials, including the Lipid Research Clinics Coronary Primary Prevention Trial ³⁸ and the Familial Atherosclerosis Treatment Study, ⁵⁶ have demonstrated the clinical benefit of bile acid sequestrants. Despite their known benefits, the use of bile acid sequestrants in clinical practice is hampered, in part because of issues related to poor palatability of the drugs and the occurrence of adverse gastrointestinal effects, particularly constipation, which influences therapy adherence. The incidence of constipation with colesevelam approximates that with placebo; with colestipol and cholestyramine, it is 10% and 26%, respectively. ^{57,58} As a result, colesevelam hydrochloride is often the preferred drug of this class.

The use of colesevelam hydrochloride in combination therapy with lovastatin, 59 simvastatin, 60 or atorvastatin 61 has demonstrated significant lowering of LDL-C levels in hypercholesterolemic patients. Low-dose combination therapy with colesevelam hydrochloride (2.3 g) and lovastatin (10 mg) was shown to reduce LDL-C by 34% (60 mg/dL; P < 0.0001) and 32% (53 mg/dL; P < 0.0001) in patients with primary hypercholesterolemia. 59 Similarly, combination therapy with simvastatin 10 mg and 20 mg and colesevelam hydrochloride 2.3 g and 3.8 g was demonstrated to reduce LDL-C. Patients treated with simvastatin-colesevelam hydrochloride combi nation therapy had a mean reduction in LDL-C levels of 42% (- 80 mg/dL; P < 0.0001) compared with baseline, which sur passed reductions for simvastatin 10 mg (- 26%, - 48 mg/dL) or 20 mg (- 34%, - 61 mg/dL) alone or for colesevelam hydro chloride 2.3 g (-8%, -17 mg/dL) or 3.8 g (-16%, -31 mg/dL) alone (P < 0.001).

Another clinical trial demonstrated that the coadministration of colesevelam hydrochloride 3.8 g and atorvastatin 10 mg or 80 mg/day resulted in LDL-C reductions of 12% to 53% in all active treatment groups (P < 0.01). Combination therapy resulted in significant decreases in LDL-C (48%) compared with colesevelam hydrochloride (12%) or low-dose atorvastatin (38%) alone (P < 0.01) but similar to those achieved with atorvastatin 80 mg/day (53%). 61

Colesevelam hydrochloride has also been demonstrated to reduce hemoglobin A1c (HbA1c) levels in subjects with type 2 diabetes mellitus uncontrolled by existing antihyperglycemic therapy (HbA1c values of 7% to 10%). The Glucose Lowering Effect of WelChol Study (GLOWS) demonstrated that 12 weeks of colesevelam 3.75 g/day was associated with significant reductions in HbA1c in 65 subjects with type 2 diabetes. 62 In subjects with a baseline HbA1c > 8.0%, the difference in least squares mean change in HbA1c was - 1.0% (0.27; P = 0.002). Compared with placebo, colesevelam treatment was associated with reductions in levels of fructosamine (-29.0 [10.9] pmol/L; P = 0.011) and postprandial glucose (- 31.5 [13.6] mg/dL; P =0.026). The mean percentage change in LDL-C was - 10% in the colesevelam group, compared with 2% in the placebo group (treatment difference, - 12% [4.2]; P = 0.007). Colesevelam was also associated with significant decreases in the percentage change in apo B (P = 0.003) and LDL particle concentration (P =0.037). 62

In another trial, colesevelam hydrochloride 3.75 g/day in subjects with type 2 diabetes mellitus uncontrolled (HbA1c 7.5% to 9.5%) with insulin, alone or in combination with oral antihyperglycemics, those receiving treatment (n = 147) experienced a significant change in HbA1c (- 0.41%) compared with controls (+ 0.09%, n = 140; P < 0.0001). Colesevelam hydrochloride also resulted in significant mean decreases in LDL-C (13%) and apo B (5%) and increases in apo AI (2%) and median triglyceride levels (21.5%). 63 The results further supported the efficacy of colesevelam hydro chloride in enhancing glycemic control and improving lipid profiles in subjects with type 2 diabetes on an insulin-containing regimen.

NIACIN

Niacin or nicotinic acid, a potent lipid-modifying agent, is a form of vitamin B $_3$, one of the water-soluble B complex vita mins. Niacin was the first lipid-lowering agent to significantly reduce cardiovascular events in the Coronary Drug Project, which randomized 3908 men and demonstrated that 6 years of niacin therapy reduced the risk of nonfatal myocardial infarction and resulted in an 11% reduction in all-cause mortality (this was not significant after 5 years of follow-up; myocardial infarction [26%] and stroke [24%] reduction were significantly compared with placebo). 64

Niacin has broad-spectrum effects including LDL-C reduction, triglyceride and lipoprotein(a) reduction, and HDL elevation and is therefore used in the treatment of a variety of lipid disorders, including the metabolic syndrome, diabetes mellitus, isolated low HDL-C, and hypertriglyceridemia. ⁶⁵ Niacin binds to the HM74 receptor on the surface of visceral adipocytes and helps reduce the mobilization of free fatty acid. Niacin inhibits intrahepatic triglyceride and VLDL biosynthesis, stimulates HDL secretion, reduces HDL catabolism, and increases LDL particle size and reduces LDL particle number. ⁶⁶

Despite its benefits in the management of dyslipidemia, the use of niacin has been limited by vasodilation-induced flushing. Niacin is currently available in three formulations, including immediate release, extended release, and long acting. Each differs with respect to its safety and efficacy profiles, with longacting or extended-release niacin having less flushing. The flushing induced by niacin is mediated by prostaglandin D $_1$. The biosynthesis of prostaglandin D $_1$ can be reduced by pretreating patients with 325 mg of aspirin (generally 2 hours before nicotinic acid is taken); limiting the ingestion of hot liquids, spicy foods, and saturated fat; and avoiding alcohol. It is recommended that over-the-counter preparations of niacin be avoided, given their side effect profile and uncertain purity. Flushing is inhibited more significantly if patients use aspirin prophylaxis at 325 mg rather than 81 mg. 67

Statin-niacin combination therapy is a recognized lipidaltering therapy used to reduce residual cardiovascular risk. Niacin is often added to a statin in patients with combined hyperlipidemia, especially if the HDL is low or lipoprotein(a) is high. Although statins have demonstrated an approximate reduction in CHD events by 30%, combination therapy with statins and niacin has resulted in relative risk reductions of ~75%. ⁶⁸ This significant reduction in CHD events suggests that other effects of niacin, such as triglyceride and lipoprotein(a) lowering and HDL raising, may also contribute to the benefits.

The HDL-Atherosclerosis Treatment Study assessed the impact of simvastatin (10 to 20 mg/day) plus niacin (2 to 4 g/day) combination therapy on risk reduction for a composite of cardiovascular endpoints (death from coronary causes, confirmed myocardial infarction or stroke, or revascularization). Cardiovascular risk was decreased by 90% in the group treated with simvastatin plus niacin compared with placebo (P = 0.03). ⁶⁹ In addition, simvastatin-niacin combination therapy resulted in a 0.4% regression in coronary stenosis, whereas progression occurred in other treatment groups receiving antioxidants alone, simvastatin plus niacin plus antioxidants, or placebo (P < 0.001). ⁶⁸ Additional studies evaluating statin plus niacin combination therapy have also demonstrated efficacy in increasing HDL-C and reducing triglycerides and LDL-C. ⁷⁰⁻⁷²

The Safety and Efficacy of A Combination of NiAcin ER and Simvastatin in PaTients (SEACOAST) study assessed the use of two combination niacin extended-release (NER) thera pies with simvastatin (1000 mg NER/20mg simvastatin and 2000 mg NER/20 mg simvastatin) in 319 dyslipidemic

patients. After 24 weeks of therapy, 6.5% and 9%, respectively, of patients discontinued use because of flushing. Similar results were observed in SEACOAST II, with combination -NER/simvastatin in higher doses (1000 mg NER/40 mg simvastatin), in which 4% and 6% (respectively) of patients withdrew because of flushing. ⁷³ An Open-Label Evaluation of the Safety and Effi cacy of a Combination of Niacin ER and Simvastatin in Patients with Dyslipidemia (OCEANS) study found comparable results with the use of NER/simvastatin (titrated to 2000/40 mg/day) during 52 weeks in 520 patients with primary type II or mixed hyperlipidemia, ⁷⁴ further substantial attainment of the benefit of statin-niacin combination therapy.

A number of trials have also demonstrated the safety of statin-niacin combination and efficacy in inhibiting the progression of atherosclerosis. Most trials have used either immediate-release niacin or extended-release niacin (Niaspan). The addition of extended-release niacin to statin therapy was evaluated in the Arterial Biology for the Investi gation of the Treatment Effects of Reducing Cholesterol (ARBITER) 2 trial, which demonstrated an increase in HDL-C by 21% and slowed progression of atherosclerosis as mea sured by change in carotid intima-media thickness, compared with statin therapy alone in patients with known CHD and low HDL-C levels. ⁷⁵

The Arterial Biology for the Investigation of the Treatment Effects of Reducing Cholesterol 6 (ARBITER 6)-HDL and LDL

Treatment Strategies in Atherosclerosis (HALTS) trial, despite some concerns about its early termination, demonstrated that in patients with CHD or CHD risk equivalent, the use of extended-release niacin 2000 mg/day in combination with long-term statin therapy caused a significant regression in carotid intima-media thickness compared with an ezetimibe-statin combination. ⁷⁶

On the basis of these surrogate outcome trials and the effect of niacin on increasing HDL-C and decreasing LDL-C and lipoprotein(a) at higher doses, niacin is a preferred treatment in combination with statin therapy to reduce cardiovascular risk. However, there is still debate as to the extent to which findings from surrogate endpoint studies can be translated into clinical benefit, and thus the completion of the clinical outcomes studies involving niacin, such as AIM-HIGH, is of great interest.

STATINS-EZETIMIBE

Combination therapy with ezetimibe, a selective cholesterol absorption inhibitor that blocks cholesterol absorption at the intestinal brush border, and statins can be used to enhance LDL-C lowering. 77,78 Ezetimibe inhibits the sterol transporter Niemann-Pick C1-like 1 protein along the luminal surface of the jejunal brush border (Fig. 14-6). Combination ezetimibe-statin therapy has been evaluated in several randomized

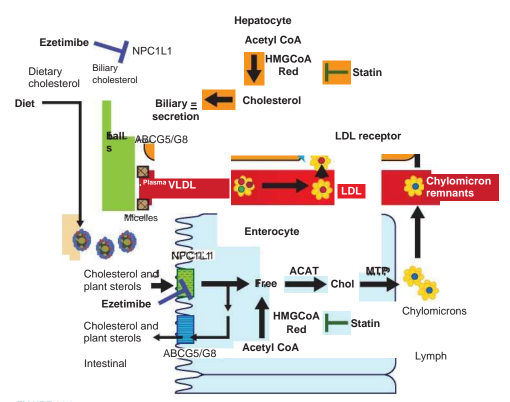


FIGURE 14-6 The mechanism of ezetimibe, a selective cholesterol absorption inhibitor. As outlined in this figure, ezetimibe reduces the small intestinal enterocyte absorption of cholesterol and plant sterols by binding to the Niemann-Pick C1-like 1 (NPC1L1) protein, which keeps cholesterol in the intestinal lumen for excretion. Excess intracellular cholesterol can also be translocated back into the intestinal lumen through the activity of the heterodimeric ATP-binding membrane cassette transport proteins G5/G8. Absorbed ezetimibe undergoes glucuronidation to a single metabolite and localizes at the intestinal wall, where it binds with higher affinity for NPC1L1 to prevent cholesterol absorption. Free cholesterol is esterified by the enzyme acyl:cholesterol acyltransferase (ACAT). Absorbed lipid, cholesteryl esters, and apo B48 are used by microsomal triglyceride transfer protein (MTP) to assemble chylomicrons. Chylomicrons are then secreted into the enteric lymphatic system and are released into the central circulation by the lymphatic duct. Chylomicrons are cleared by the liver. Ezetimibe can also reduce the hepatic reabsorption of cholesterol from bile by inhibiting NPC1L1 along the canalicular surface of the biliary tract. (From Davis HR, Veltri EP: Zetia: inhibition of Niemann-Pick C1 like 1 [NPC1L1] to reduce intestinal cholesterol absorption and treat hyperlipidemia. J Atheroscler Thromb 14:99, 2007. Reproduced with permission.)

clinical trials of patients with primary hypercholesterolemia. Ezetimibe 10 mg coadministered with simvastatin 10 mg resulted in 44% LDL-C reductions, similar to those obtained with simvastatin 80 mg alone, and coadministration of ezetimibe 10 mg with simvastatin doses of 10 to 80 mg resulted in triglyceride reductions of 26% to 31% and HDL-C increases of 8% to 11%. ⁷⁹ In another study, coadministration of ezeti mibe 10 mg and atorvastatin 10, 20, 40, or 80 mg compared with monotherapy resulted in significant improvements with a 12% reduction in LDL-C, 8% reduction in triglycerides, and 3% increase in HDL-C. ⁸⁰

Lovastatin monotherapy at 10, 20, or 40 mg or ezetimibe 10 mg was compared with combination therapy of ezetimibe 10 mg plus lovastatin 10, 20, or 40 mg; the coadministration of ezetimibe provided an incremental 14% decrease in LDL-C, a 5% increase in HDL-C, and a 10% decrease in triglycerides compared with pooled lovastatin alone. Ezetimibe plus lov astatin provided mean LDL-C decreases of 33% to 45%, median triglyceride decreases of 19% to 27%, and mean HDL-C increases of 8% to 9%, depending on the statin dose. ⁸¹

In clinical practice, the prevailing recommendation is to initiate LDL-C reduction with statin therapy. If the patient cannot attain the LDL-C goal with statin monotherapy either because of an inadequate response or because of intolerance to an appropriate dose of a statin, then either niacin, a resin, or ezetimibe can be added as adjuvant therapy. Ezetimibe monotherapy can also be used in patients who are intolerant to statin therapy. For patients receiving statin therapy not at goal, the dose of the statin should generally be titrated to at least 40 mg/day before ezetimibe is added. For patients who are more than 10% above their LDL-C goal, statin titration may result in a 6% further LDL-C reduction, and ezetimibe may therefore provide enhanced ability to achieve LDL-C targets in these patients.

Ezetimibe therapy provides a level of LDL-C reduction that is equivalent to three titration steps of a statin. The efficacy of ezetimibe for reducing cardiovascular morbidity and mortality in patients with CAD is being evaluated in the IMProved Reduction of Outcomes: Vytorin Efficacy International (IMPROVE-IT) trial. 82 Unfortunately, we will probably not have clinical outcome results from this pivotal clinical trial until 2013 at the earliest.

STATIN-FIBRATE COMBINATION THERAPY

In patients with combined dyslipidemia, fibrate-statin combination therapy can be used to promote reductions in LDL-C and triglycerides and simultaneous increases in HDL-C. The fibrates activate lipoprotein lipase and promote triglyceride hydrolysis, inhibit hepatic triglyceride biosynthesis, stimulate hepatic HDL secretion, and reduce LDL particle number. Fibrate monotherapy has been shown to reduce the risk for cardiovascular events ^{83,84} and rates of atherosclerotic disease progression. ^{85,86}

The Simvastatin Plus Fenofibrate for Combined Hyperlip - idemia (SAFARI) trial demonstrated that combination therapy of simvastatin 20 mg/day plus fenofibrate 160 mg/day resulted in significantly decreased LDL-C levels (31%) com pared with monotherapy (26%; P < 0.001) in patients with combined hyperlipidemia (fasting triglyceride levels > 150 and < 500 mg/dL and LDL cholesterol > 130 mg/dL). ⁸⁷ In addition , mean HDL-C levels significantly increased with combination therapy (19%) compared with monotherapy (10%; P < 0.001), without the occurrence of any drug-related serious adverse events. ⁸⁷

Fibrates are associated with a slightly increased risk (<1.0%) for myopathy, cholelithiasis, and venous thrombosis. When fibrate therapy is used, measurement of serum creatinine -concentration is recommended before fibrate use, with dose adjustments made for renal impairment. Whereas routine monitoring of creatinine concentration is not required, a

clinically important increase in creatinine without other potential causes should be reevaluated, and consideration should be given to discontinuation of fibrate therapy or reduction of the dose. ⁸⁸ The efficacy of fenofibrate used in combination with simvastatin compared with simvastatin monotherapy is the Action to Control Cardiovascular Risk in Diabetes trial. Although the primary composite endpoint reduction between the two groups as a whole was not significantly different, among subjects with elevated triglycerides (> 204 mg/dL) and low HDL (< 34 mg/dL), there was a trend for improved benefit with combination therapy (31%). ^{88a}

STATIN-OMEGA-3 FATTY ACIDS

Omega-3 fatty acids, or fish oils, significantly reduce VLDL and triglyceride levels and increase LDL-C levels in patients with high triglycerides. The omega-3 fatty acids inhibit the enzyme diacylglycerol acyltransferase 2, thereby reducing intrahepatic triglyceride biosynthesis. They also stimulate mitochondrial beta-oxidation of fatty acids, decrease VLDL production and biosynthesis, and stimulate triglyceride hydrolysis by lipoprotein lipase. Dietary supplementation with the n-3 polyunsaturated fatty acids (PUFAs) eicosapen taenoic acid (EPA) and docosahexaenoic acid (DHA) has also been shown to lower the risk of death, nonfatal coronary events, and stroke after myocardial infarction. ⁸⁹ In several clinical trials, PUFAs have been shown to reduce triglyce ride levels by 20% to 30% and by up to 50% in patients with severe hypertriglyceridemia (triglycerides > 500 mg/dL [> 5.65 mmol/L]). ^{90,91}

Combination therapy with statins and n-3 PUFAs has demonstrated LDL-C reductions of 13% to 24% and triglyceride reductions of 27% to 30% added to pravastatin 40 mg/day 92 or simvastatin 20 mg/day. 93 Similarly, combination therapy with atorvastatin 10 mg resulted in significant reductions of the concentration of small, dense LDL particles and increases in HDL-C compared with monotherapy. 94 In a study evaluated in the effects of adding prescription omega-3 acid ethyl esters 4 g/day (P-OM3; Lovaza, formerly Omacor) to simvas tatin 40 mg in more than 250 patients with persistent hyper triglyceridemia, decreases in non-HDL-C were significantly greater with combination therapy than with placebo plus simvastatin (9.0% versus 2.2%, respectively; P < 0.001). In addition, combination therapy significantly lowered triglyc erides (29.5%) and VLDL-C (27.5%), raised HDL-C (3.4%), and lowered the ratio of total cholesterol to HDL-C (9.6%; P < 0.001 versus placebo for all). 95

The Japan EPA Lipid Intervention Study (JELIS) evaluated whether the addition of fish oils to patients already taking a statin would provide incremental risk reduction. Approximately 19,000 Japanese men and women with hypercholes terolemia were prospectively randomized to statin therapy with or without 1800 mg/day of EPA. ⁹⁶ Combination therapy resulted in an additional 19% reduction in major coronary events at 4.6 years of follow-up compared with statin monotherapy.

BEYOND LDL-C

Whereas LDL-C has become the cornerstone of lipid diagnosis and therapy, apo B and non-HDL-C may be more accurate indicators of cardiovascular risk as well as measures by which to gauge LDL-lowering therapy. ⁹⁷ The measurement of apo B provides an estimate of the number of atherosclerotic particles, and non-HDL-C represents the sum of cholesterol in VLDL and LDL, or the mass of cholesterol and cholesteryl

FIGURE 14-7 Estimated change in relative risk reduction for nonfatal myocardial infarction (MI) and coronary heart disease (CHD) death associated with mean LDL-C reduction. The 95% probability interval is represented by the dotted line. The crude risk estimates from individual statin trials are plotted with their 95% confidence intervals. (From Ridker PM, Danielson E, Fonseca FA, et al: Reduction in C-reactive protein and LDL cholesterol and cardiovascular event rates after initiation of rosuvastatin: a prospective study of the JUPITER trial. Lancet 373:1175, 2009; Robinson JG, Smith B, Maheshwari N, Schrott H: Pleiotropic effects of statins: benefit beyond cholesterol reduction? J Am Coll Cardiol 46:1855, 2005. Reproduced with permission.)

ester within apo B particles. ⁹⁷ Defined more specifically, non-HDL-C (total cholesterol minus HDL-C) reflects the concentration of cholesterol within all lipoprotein particles currently considered atherogenic. ⁹⁸ Whereas debate continues as to whether apo B or non-HDL-C directed therapy may be more effective, it is evident that both have advantages over LDL-C as predictors of cardiovascular risk.

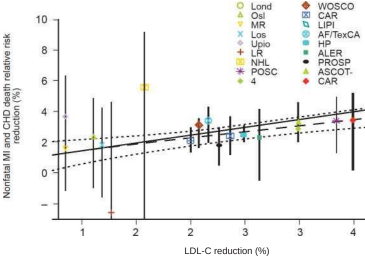
In several epidemiological studies and in post hoc analyzes of clinical trials, apo B has been found to be a better predictor than LDL-C of CVD risk. ^{99,100} Additional analyzes suggest that once LDL-C is lowered, apo B may be a more effective way to assess residual CVD risk and to determine the need for medication adjustments. ⁹⁸ A recent consensus panel building on the guidelines of the NCEP ATP III and the American Heart Association/Centers for Disease Control for cardiovascular risk assessment recommended that for patients with diabetes mellitus and one additional risk factor, the goals of treatment should be LDL-C < 70 mg/dL and non-HDL-C < 100 mg/dL, with optional targets of apo B < 80 mg/dL or LDL particle number (LDL-P) < 1000 nmol/L. ¹⁰¹

COMPLIANCE WITH MEDICATIONS

Adherence to LDL-lowering therapy is a key component in achieving target goals. A Canadian cohort of 85,020 primary prevention patients showed that after 6 and 24 months, the percentage of patients remaining on statin therapy for dyslipidemia treatment was 50% and 24%, respectively. 102 In a managed care population of high-risk patients, better compliance with statin therapy was associated with older age, being male, more frequent outpatient follow-up for lipid management, history of hospitalization for a coronary event, history of revascularization, and receiving prescriptions through the mail rather than having to pick them up monthly at a retail pharmacy. 103 The NCEP guidelines identify that adherence issues need to be addressed for the highest possible levels of CHD risk reduction to be achieved. Patient education and involvement in self-monitoring, provider education and reinforcement of lipid treatment guidelines, and use of health delivery system strategies (including lipid management clinics, collaborative care with pharmacists, the use of critical care pathways, cardiac rehabilitation for patients who sustain acute cardiac events or who undergo revascularization, tele health technologies, and nursing case management) are identified as targets for intervention to improve adherence. 31 The need for compliance with lipid-lowering medications should be reinforced at each office visit.

CONCLUSION

The strength of LDL-C as a target for CVD risk reduction is supported by scientific evidence based on the biology of atherosclerosis, epidemiology, and clinical trial evidence with both statin- and non-statin-based approaches to serum lipid lowering. A complex set of biochemical reactions promotes the development of inflammation, LDL oxidation, and plaque progression. The absence of LDL-C elevation and other atherogenic lipoproteins is associated with less atherosclerosis in



animal models. Life-long low LDL-C levels (as seen in PCSK9 loss-of-function mutations) in humans are associated with very low rates of CVD. Clinical trials using statins and non-statin approaches to lipid lowering have demonstrated that for every 1 mg/dL reduction in serum LDL-C, there is a 1% reduction in risk for acute cardiovascular events. A number of clinical trials have confirmed that LDL-C lowering predicts CVD risk reduction (Fig. 14-7). 104,105 The totality of the evidence strongly supports that LDL-C and, more recently, non-HDL-C are targets that, if achieved, result in a marked reduction in CVD. The challenge for clinicians is to identify the most appropriate patients for LDL-C or non-HDL-C treatment and maintain adherence to therapy to improve outcomes and quality of life.

On-target LDL-C predicts the benefit of treatment, and therapeutic lifestyle changes coupled with statin therapy remain the mainstay of treatment. Combination therapy has demonstrated greater benefit with lipid management, not just in terms of LDL-C but also with respect to HDL-C and triglycerides. During the next several years, ongoing clinical trials will evaluate the impact of comprehensive lipid management (modifying triglycerides, HDL-C, and other lipid parameters) compared with intensive LDL-C lowering alone and may provide further guidance on strategies to optimize cardiovascular risk reduction.

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CHAPTER 15

The Contribution of Triglycerides and **Triglyceride-Rich Lipoproteins to** Atherosclerotic Cardiovascular Disease

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KEY POINTS

- · When TG levels are between 200 and 800 mg/dL, TG-rich particles are associated with the presence of small dense LDL, low levels of HDL-C. insulin resistance, and metabolic syndrome, all of which increase risk for atherosclerosis.
- TG-rich particles affect the size and atherogenic potential of VLDL remnants and LDL particles, both of which enter the arterial subendothelial space and contribute to atherosclerotic plaque.
- In a meta-analysis of 68 prospective studies, TG level was not a predictor of risk for nonfatal myocardial infarction and death after adjustment whereas non-HDL-C was associated with increased risk after adjustment for HDL and log
- Non-HDL-C predicted risk of CHD and future cardiovascular events better than LDL-C did, probably because non-HDL-C includes all apo B-containing lipoproteins.
- Nonfasting TG levels may predict CHD better than fasting levels but are more difficult to standardize and to measure in the clinical setting; non-HDL-C can be measured in the nonfasting state.
- NCEP ATP III recommends optimal TG level <150 mg/dL. NCEP recommends calculation of non-HDL-C when TG >200 mg/dL, with the goal being 30 mg/dL higher than the LDL-C goal.

- · Lifestyle changes-exercise, diet, and weight lossare necessary to lower TG levels, especially in patients with metabolic syndrome.
- · Statins, fibrates, niacin, and omega-3 fatty acids can be used to lower TG levels if lifestyle changes are insufficient to reach the goal.

The contribution of triglyceride (TG) and TGrich lipoproteins to the development of atherosclerosis, especially as an independent predictor of cardiovascular risk, has been debated for many years. TG levels between 200 and 800 mg/dL may be associated with other lipid abnormalities that predispose to atherosclerosis, including low levels of highdensity lipoprotein choles terol (HDL-C), 1-3 small, dense low-density lipoprotein (LDL) particles, atherogenic TG-rich remnants, 4,5 and insulin resistance, 6,7 all of which increase the risk for coronary heart disease (CHD). Therefore, it is difficult to determine the independent contribution of TG and how should aggressively one hypertriglyceridemia. The percentage of adults in the United States with TG levels above 150 mg/dL (1.7 mmol/L), 200 mg/dL (2.3 mmol/L), 500 mg/dL (5.7 mmol/L), and 1000 mg/dL (11.3 mmol/L) is 33%, 18%, 1.7%, and 0.4%, respectively. 8 Therefore, hypertriglyceridemia affects a significant portion of the population. Reliable assessment of the risk associated with lipid fractions is important for the development of accurate screening and treatment strategies.

This chapter reviews the epidemiological evidence indicating that hypertriglyceridemia contributes atherosclerosis. metabolism is then described to demonstrate the mechanisms by which the level of TG-rich lipoproteins affects the composition and size of remnant, LDL, and HDL particles and how this affects the development of atherosclerosis and risk for CHD. Finally, the classification of various types causes hypertriglyceridemia are discussed, followed by suggestions for evaluation, treatment, and management.

EPIDEMIOLOGIC EVIDENCE LINKING TRIGLYCERIDE LEVELS WITH RISK FOR CORONARY **HEART DISEASE**

epidemiological studies Several provided important findings on the role of TG-rich lipoproteins and atherosclerosis. In 1959, Albrink and Man 9 observed that TG levels greater than 175 mg/dL were present in 70% of 100 cases of myocardial infarction (MI) compared with only 7% of 92 healthy controls. In the 8-year Cardiovascular Prospective (PROCAM) study reported in 1996, elevated levels of TG were independently associated with incident CHD events after adjustment for LDL-C and HDL-C levels. 10 For this reason, elevated TG levels are in the European model for calculation of cardiovascular risk. 11 An important contribution of PROCAM was the observation that an increasing TG level is directly associated with CHD incidence up to a level of 800 mg/dL. Levels higher than 800 mg/dL are thought not to be associated with CHD because of lipoprotein particles too large to penetrate the vascular endothelium compared with smaller, atherogenic remnant particles found with mild hypertriglyceridaemia. In the Framingham Heart Off spring Study, TG level was not associated with CHD after adjustment for HDL-C and other covariates in both men and women. 12 However, a later analysis showed that the cholesterol in remnant lipoproteins is an independent predictor of CHD risk in Framingham women. 13

In 1996, the first large meta-analysis of 17 prospective, observational studies of TG and CHD events reported that an increase of 89 mg/dL (1 mmol/L) in TG level was

associated with a univariate risk of 32% in men and 76% in women and was an independent risk predictor (adjusted for other covariates including HDL-C) of 14% in men and 37% in women. ¹⁴

The Baltimore Coronary Observational Long-Term Study (COLTS), a retrospective cohort study, observed 350 individuals with CHD for up to 18 years after cardiac catheterization . After adjustment for age, gender, use of beta blockers, and other risk factors, baseline fasting TG level > 100 mg/dL was associated with a 50% increased risk of subsequent events compared with those with TG < 100 mg/dL (P = 0.008) and was an independent predictor of recurrent cardiovascular events in patients with CHD. ¹⁵ These results suggested that an optimal TG level is < 100 mg/dL in patients with CHD.

The Copenhagen Male Study was a prospective study of 2906 white men without CHD at baseline. ¹⁶ During 8-year follow-up, there was an increased risk for CHD with increasing TG tertiles: 4.65% for TG of 0.4 to 1.1 mmol/L; 7.7% for TG of 1.1 to 1.6 mmol/L; and 11.5% for TG of 1.6 to 2.2 mmol/L. After adjustment for age, body mass index, hypertension, smoking, alcohol, physical activity, diabetes, socioeconomic status, LDL-C, and HDL-C, the middle and highest level of TGs had a relative risk of 1.5 and 2.2, respectively, compared with the lowest tertile of TGs. Moreover, elevated TG and low HDL-C levels were predictive of CHD events in men with LDL-C both less than and greater than 170 mg/dL. ¹⁷

In 2004, a meta-analysis pooled individual data from 96,224 subjects in 26 prospective studies in New Zealand, Australia, and several Asian countries and included 670 and 667 deaths from CHD and stroke, respectively. ¹⁸ After adjustment for age, sex, blood pressure, smoking, ratio of total cholesterol (TC) to HDL-C, and major cardiovascular risk factors, compared with those in the bottom fifth of TG levels, individuals in the top fifth of TG levels had a 70% (95% CI, 47-96) greater risk of CHD death, an 80% (95% CI, 49-119) higher risk of fatal or nonfatal CHD, and a 50% (95% CI, 29-76) increased risk of fatal or nonfatal stroke. The association between levels of TG and CHD death was similar across subgroups defined by ethnicity, age, and sex. These results suggested that serum TGs are an independent predictor of CHD and stroke risk in the Asia-Pacific region even after adjustment for HDL-C.

More recently, a meta-analysis of 29 population-based, Western prospective studies included 10,158 CHD cases from 262,525 participants. After correction of risk estimates for longterm within-individual variation in TG measurements, the risk of CHD adjusted for age, gender, and calendar period was approximately twofold higher in individuals in the top third of log TG levels compared with the bottom third. ¹⁹ The attributable risk was similar in men and women and in fasted and non-fasted participants. After adjustment for HDL-C, the odds ratio for CHD was attenuated to 1.72 but remained sig nificant (95% CI, 1.56-1.90) in those in the top third of log TG level compared with the bottom third. An important contribution was that repeated measurements an average of 4 years apart in 1933 participants in the EPIC-Norfolk study and an average of 12 years apart in 379 participants in the Reykjavik study showed that the long-term stability of log TG values is similar to that of blood pressure and total serum cholesterol (within-person correlation coefficients of 0.64 [95% CI, 0.60-0.68] during 4 years and 0.63 [95% CI, 0.57 0.70] during 12 years). 20

The Metabolic, Lifestyle and Nutrition Assessment in Young Adults (MELANY) study obtained two measurements of fasting TGs 5 years apart, as well as lifestyle variables, in 13,953 healthy men, aged 26 to 45 years, who were then observed for an average of 10.5 years. ²¹ After adjustment for eating breakfast, smoking, exercise, and changes in body mass index, those in the top quintile of TGs had a fourfold higher risk of angiographically proven CHD compared with those in the lowest quintile. The magnitude of this risk was much greater than the average 1.7-fold increase in risk observed in the 2007 meta-analysis ¹⁹ and

other large studies, a finding thought secondary to the younger age of the cohort compared with other studies. Another important finding was that change in TG level between the initial and the second measurements was positively associated with change in CHD risk.

In a recent subgroup analysis of the Treating to New Targets (TNT) study and Incremental Decrease in Endpoints through Aggressive Lipid Lowering (IDEAL) study, the utility of TGs to predict new cardiovascular events was examined. 22 IDEAL compared atorvastatin 80 mg with simvastatin 20 to 40 mg and TNT compared atorvastatin 80 mg with atorvastatin 10 mg in patients with CHD or a history of MI. After adjustment for age and gender, the risk of cardiovascular events was 63% higher in patients in the highest quintile of TG (HR, 1.63; 95% CI, 1.46-1.81) compared with the lowest quintile. The ability of TGs to predict risk was attenuated when HDL and apo B/apo AI were in the model, and it was eliminated with inclusion of diabetes, body mass index, glucose, hypertension, and smoking (P = 0.044 and 0.621, respectively, for the trend across quintiles of TG). Similar results were observed in those in whom LDL-C had been lowered to goal.

The Pravastatin or Atorvastatin Evaluation and Infection Therapy-Thrombolysis in Myocardial Infarction (PROVE IT-TIMI 22) trial randomized patients with acute coronary syndromes to atorvastatin 80 mg or pravastatin 40 mg. ²³ An ontreatment TG level < 150 mg/dL was independently associated with a significant reduction in risk for the composite primary endpoint of nonfatal MI, death, and recurrent acute coronary syndrome. After adjustment for LDL-C and other covariates in a subanalysis, each 10 mg/dL decline in on-treatment TG level was associated with a 1.6% lower risk of the primary endpoint.

In summary, several studies show TGs to be an independent predictor of CHD after adjustment, whereas in others, risk of TGs is attenuated or eliminated after adjustment, especially for HDL-C levels. In the next section, potential reasons for these findings are examined on the basis of the metabolic interrelationships between levels of TG-rich lipoproteins, small dense LDL, remnant particles, and HDL particles.

LIPOPROTEIN METABOLISM

Determination of the contribution of TGs to risk for atherosclerosis is complicated by the fact that the metabolism of TG-rich lipoprotein fractions significantly affects the levels and composition of other lipoprotein fractions that also contribute to cardiovascular risk. ²⁴Therefore, in evaluating the role of TG and TG-rich lipoproteins in contributing to atherosclerosis, one must consider the complex interrelationships between the various lipoproteins during metabolism.

TGs (triacylglycerols) are composed of a glycerol backbone in which each of the three hydroxyl groups is esterified with a fatty acid. 25,26 TGs play an important role in lipid metabolism and are the major source of metabolic energy storage. Cholesterol and TGs are almost insoluble in plasma; there fore, they are transported in lipoprotein particles from the liver (endogenous production of very-low-density lipoprotein [VLDL]) and intestine (exogenous production of chylomi crons [CMs] from dietary fat) to various tissues-TGs for energy use in skeletal muscle or storage in adipose tissue and cholesterol for synthesis of steroid hormones, bile acid formation, and cell membrane structural integrity. Lipoprotein particles are spherical and contain a central core of varying amounts of TG and cholesteryl ester (CE) (both nonpolar lipids) covered on the surface by a monolayer of polar lipids (primarily phospholipids), one or more apolipoproteins, and



Apo B exists in two forms in plasma, apo B100 and apo B48, both of which are products of the same structural gene on chromosome 2. ²⁷ Both apo B48 and apo B100 are constitutively synthesized; the availability of TG and CE (the core lipids) regulates their secretion. Containing mainly TG in their core, CMs and VLDL are the major TG carriers in plasma and are the two largest classes of lipoproteins. There are two pathways for the metabolism of TG-rich lipoproteins; the exogenous pathway carries dietary fats by apo B48 in CMs, whereas the endogenous pathway represents hepatic secretion of VLDL, a TG-rich apo B100-containing lipoprotein.

Endogenous Pathway: Assembly of VLDL Apo B-100 Lipoprotein Particles

The full-length apo B100 is a glycoprotein that contains 4536 amino acids. Apo B100 is synthesized by the liver and secreted in the form of VLDL, a TG-rich-lipoprotein that in plasma contains 60% TG by mass and 20% CE by mass. 27 Both fatty acids synthesized de novo from acetyl coenzyme A and fatty acids from lipolysis of stored adipose tissue TG or from core lipids of TG-rich remnant particles returning to the liver stimulate the assembly of VLDL in the liver (Fig. 15-1). Microsomal triglyceride transfer protein (MTP) transfers TGs from the cytosol to the endoplasmic reticulum containing nascent apo B during the assembly of CM and VLDL in enterocytes and hepatocytes, respectively. ²⁸ MTP gene expression is regulated by insulin, possibly through transcriptional activity of the sterol response element-binding protein 1c (SREBP-1c). This may explain why VLDL secretion and TG levels are increased in insulin resistance syndromes. ²⁹ In the plasma (see Fig. 15-1), VLDL particles adhere to gly cosaminoglycan molecules on endothelial cells of capillaries, primarily in muscle, lung, and adipose tissue, 30,31 where interaction with lipoprotein lipase (LPL) and glycosylphosphati dylinositol-anchored high-density lipoprotein-binding protein 1 (GPIHBP1) results in hydrolysis of VLDL triglycerides to free fatty acids and glycerol. 32,33

The removal of TG from VLDL by LPL exposes the apo E molecules on the lipoprotein surface of VLDL. ³² Apo E functions as a ligand in the receptor-mediated clearance of CM and VLDL remnants in the liver through several receptors: a remnant receptor, 34-36 the LDL receptor (an apo B/apo E receptor), the LDL receptor-related protein, the VLDL receptor, and apo E receptors. 37 With stable isotope methodology, Welty and colleagues 38 published the first study of simultaneous ous kinetics of apo B48 and apo B100 in human subjects and showed that about 50% of VLDL is directly removed from plasma (see Fig. 15-1) and therefore not converted to IDL or LDL. During the removal of the TG from the remaining 50% of VLDL (referred to as the delipidation cascade), the VLDL particles are hydrolyzed by LPL to smaller particles termed VLDL remnants or IDL (relatively enriched in CE but also containing TG), which then interact with hepatic lipase (HL) and are converted to CE-rich LDL, the major cholesterol-carrying lipoprotein in normal human plasma. Apo B100 is the main structural protein of LDL and contains the LDL receptor-binding domain; therefore, LDL is removed from the circulation by binding mainly to hepatic LDL receptors (see Fig. 15-1). ²⁷

Exogenous Pathway: Assembly of Chylomicrons

Produced in the intestine in response to dietary fat, CMs contain apo B48, the amino-terminal 48% of apo B100, which is synthesized by the intestine and produced by a premature stop codon at the apo B100 codon 2153 by tissue-specific mRNA processing (see Fig. 15-1). ³⁹ Genes related to sterol absorption (ABCG5) ^{40,41} and nuclear receptors LXR and FXR ⁴² in bile and sterol metabolism may affect variability in absorption of dietary fats and CM formation. Within the intestinal cell, free fatty acids combine with glycerol to form TGs, and cholesterol is esterified by acyl coenzyme A:cholesterol acyltransferase (ACAT) to form cholesterol esters.

CMs enter lacteals in the intestinal villi and travel through the lymphatics to the thoracic duct and then into the blood stream. Similar to the metabolism of VLDL, CMs bind to LPL on the surface of endothelial cells, where most of the TGs and some surface glycerophospholipids are catabolized to form CM remnants and free fatty acid by LPL, present in capillary walls primarily of skeletal muscle, adipose tissue, and lung, with apo C-II as a cofactor and apo C-III as an inhibitor (see Fig. 15-1). Apo B48 does not contain an LDL receptor binding domain; therefore, the CM remnants are taken up by the liver by remnant receptors ⁴³⁻⁴⁶ as well as by the LDL receptor that recognizes apo E. ^{47,48} CMs also bind to HL on the sinusoidal surface of hepatocytes, where HL further hydrolyzes remnant lipids. In the fed state, both CMs and VLDL transport TGs to peripheral organs, where, through the action of tissue-specific LPL, TG-derived free fatty acids are used as energy in muscle and in other tissues, converted to TG, or stored in adipose tissue. 49 Residual TGs (in the form of free fatty acids) and dietary cholesterol are rerouted to hepatocytes.

To transport TG and cholesterol to peripheral tissues, lipo protein particles cross the endothelial barrier in blood vessels to reach the extracellular space (see Fig. 15-1). The subendothelial retention of apo B100-containing lipoproteins by a chargemediated interaction with proteoglycans in the extra cellular matrix is thought to be the initiating event in atherogenesis. Smaller, electronegative LDL particles penetrate the endothelial barrier 1.7-fold better than large LDL particles do; CM and VLDL remnants can also cross. All of these particles interact with positively charged intimate proteoglycans. 50,51 Oxidation, by reactive oxygen species, of fatty acids of surface phospholipids of the apo B-containing lipoprotein particles results in modification of lysine residues of apo B. Scavenger receptors on macrophages recognize modified apo B, and unregulated uptake of the modified lipoprotein particles causes macrophage accumulation of lipids, a process leading to a foamy cytoplasm and the term foam cells (see Fig. 15-1). 52-54 The most important scavenger receptor is CD36 (also called scavenger receptor B). 55-⁵⁷ As foam cells increase in number, the fatty streak develops. VLDL particles from patients with hypertriglyceridemia are enriched in apo E, which can lead to a conformational change in the VLDL particles that facilitates binding to the macrophage scavenger receptor, resulting in unregulated uptake similar to that seen with oxidized LDL. 58 CM remnants are also small enough to enter the subendothelial space, where they are taken up by macrophages that promote atherogenesis. 59,60 Similar to LDL, IDL is taken up by macrophages and can also cause foam cell formation; endothelium-dependent vasomotor function in human coronary arteries is impaired by both IDL and LDL. 61 Several angiographic trials of cholesterol-lowering therapy



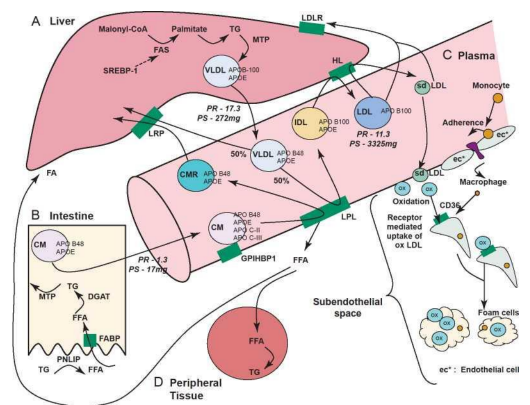


FIGURE 15-1 Schematic overview of TG-rich lipoprotein metabolism in the nonfasting (fed) state. The main sources of plasma TG are exogenous, from dietary fat, and endogenous, from the liver. In the intestine, dietary TG is hydrolyzed by pancreatic lipase (PNLIP) into monoglycerides and free fatty acids (FFA), forming micelles. Fatty acid-binding protein (FABP) transports FFA from the intestinal lumen into enterocytes, and TG is resynthesized through the sequential acyltransferase reactions, of which microsomal diacylglycerol acyltransferase (DGAT) catalyzes the terminal committed step. Microsomal triglyceride transfer protein (MTP) mediates assembly of TG with apo B48 and apo E into chylomicrons. After secretion into the blood through the lymph, chylomicrons acquire apo C-II and C-III, which modulate the plasma metabolism of TG-rich lipoproteins. Apo C-II is an obligatory cofactor for lipoprotein lipase (LPL), whereas apo C-III may interfere with LPL. Apo AV has been shown to enhance hydrolysis of TG-rich lipoproteins, through an incompletely defined mechanism; a newly characterized protein, glycosylphosphatidylinositol-anchored high-density lipoprotein-binding protein 1 (GPIHBP1), appears to facilitate lipolysis by anchoring chylomicrons to endothelium, providing stability for LPL activity. Apo B48 is the signature protein of intestinally derived TG-rich lipoproteins. In the liver, TG is synthesized from FFA extracted from plasma, from fatty acids synthesized de novo, and from uptake of plasma lipoproteins. A central enzyme for de novo synthesis is fatty acid synthase (FAS), which catalyzes the conversion of malonyl-CoA to palmitate. FAS is induced by the membrane-bound transcription factor sterol regulatory element-binding protein 1 (SREBP-1), which is itself regulated by polyunsaturated fatty acids, glucose, and insulin. Hepatic MTP mediates TG assembly with cholesteryl esters, apo B100, and apo E to form VLDL, which is released into the space of Disse by exocytosis. Apo B48 is the signature protein of intestinally derived TG-rich lipoproteins. In adipose and muscle capillaries, TG in chylomicrons and VLDL is hydrolyzed into FFA by endothelium-bound LPL. FFA is then re-esterified and stored in adipocytes or oxidized for energy in myocytes. Chylomicrons and VLDL are remodeled, respectively, into the smaller, denser, cholesteryl ester (CE)enriched chylomicron remnants (CMR) and VLDL remnants (also called intermediate-density lipoprotein [IDL]). CMR and some IDL are cleared by apo E-mediated endocytosis through hepatic remnant receptors (LRP). IDL can also be hydrolyzed by hepatic lipase (HL), making smaller, CE-rich LDL particles: 50% of VLDL is directly removed from plasma in normalipidemic human subjects, Production rate (PR), pool size (PS), and fractional catabolic rate for apo B48, VLDL apo B100, and LDL apo B100 are shown on a 36% fat diet. PR refers to secretion rate for VLDL apo B100 from the liver and apo B48 from the intestine. (Modified from Hegele RA, Pollex RL: Hypertriglyceridemia: phenomics and genomics. Mol Cell Biochem 326:36, 2009.)

have shown that serum IDL concentrations are predictive of an increased incidence of CHD62 and an increased incidence of coronary events in those with CHD, independently of other factors.63-65 VLDL and IDL have been identified in human atherosclerotic plaques,66 and these particles are associated with progression of mild to moderate coronary lesions. In the Monitored Atherosclerosis Regression Study (MARS) angiographic trial, IDL, but not VLDL or LDL, was associated with progression of carotid artery intima-media thickness.67 Moreover, the total TG level and markers for TG metabolism predicted risk of progression of low-grade but not of high-grade coronary artery lesions.68,69

Effect of Triglyceride-Rich Lipoproteins on Level of HDL-C in Humans

Levels of TG-rich lipoproteins also affect HDL level and particle size. HDL levels are inversely related to risk of CHD. 70-74 In contrast to LDL and VLDL, HDL has antiatherogenic properties that include reverse cholesterol transport, antioxidation (protecting apo B lipoproteins from oxidation), antithrombotic and anti-inflammatory properties, and maintenance of endothelial function. Apo A-I is secreted from the liver in a lipid-poor form. The major formation of HDL particles occurs when lipid-poor apo A-I interacts with the ABCA1

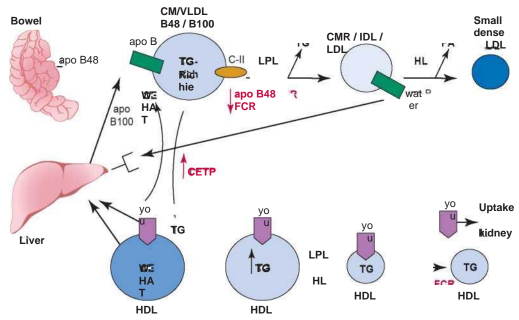


FIGURE 15-2 HDL particles play a significant role in the delivery of cholesterol from peripheral cells to the liver, a process known as reverse cholesterol transport. There are two pathways by which this can occur. In the first, the scavenger receptor class B type 1 mediates hepatic uptake of cholesterol, in the form of cholesteryl esters, from HDL particles without uptake of the whole particle or apo AI. In the second pathway, cholesteryl ester transfer protein (CETP) catalyzes the transfer of CEs from HDL-containing lipoproteins to apo B-containing lipoproteins (VLDL and CM) in exchange for triglyceride from the apo B-containing lipoproteins to HDL-containing lipoproteins. Apo AI fractional catabolic rate (FCR) is inversely correlated with the FCR of apo B48 (r = .48; P = .05) but not with VLDL apo B100 FCR or production rate. ⁸¹ Thus, when chylomicron apo B48 clearance is delayed (represented by decreased apo B48 FCR in the figure), TG-rich apo B particles accumulate and the TG is transferred to HDL apo AI particles in exchange for CE. These TG-enriched HDL particles are remodeled by hepatic lipase as well as by endothelial lipase to small TG-rich, CE-depleted HDL particles that are catabolized faster than large, CE-rich HDL; thus, the apo AI FCR is increased (represented by increased apo AI FCR in the figure), a finding resulting in lower levels of HDL-C in the setting of high TG levels. Hepatic lipase then hydrolyzes the TGs within the TG-rich LDL to release free fatty acids, a process that remodels the LDL particles into smaller and more dense LDL particles that can enter the arterial intima more easily than larger LDL particles, thus making them more atherogenic.

receptor on the surface of peripheral cells. ²⁵ This results in transfer of free cholesterol and phospholipid to the apo AI, forming a pre- \$\\$ particle. When the free cholesterol is esterified under the action of lecithin-cholesterol acyltransferase (LCAT), this particle becomes a mature HDL particle. The majority of the proteins of HDL, apo AI, A-II, and A-IV, are secreted as components of VLDL, which are then transferred to the apo AI particle in the plasma. Plasma HDL is mainly assembled extracellularly during the transfer of surface components of TGrich lipoproteins, including phospholipids and cholesterol.

HDL particles play a significant role in delivering cholesterol from peripheral cells to the liver after esterification within the particle to CE through plasma LCAT, a process known as reverse cholesterol transport. There are two paths by which this can occur. In the first, the scavenger receptor class B type 1 (SR-B1) mediates hepatic uptake of CE from HDL particles without uptake of apo AI or the whole HDL particle. 75 In the second pathway, cholesteryl ester transfer protein (CETP) catalyzes the transfer of CE from HDL to apo B-containing lipoproteins (VLDL and LDL) in exchange for TG from the apo B-containing lipoproteins (Fig. 15 - 2). 25,76,77 This exchange results in apo Bcontaining lipoproteins, which are enriched with CEs and depleted of TGs, and HDL particles, which are depleted of CEs and enriched with TGs. The TG-rich and CE-poor HDL particles are catabolized faster than the large, CE-rich HDL particles are 78 (apo AI fractional catabolic rate [FCR] is increased as noted in Fig. 15-2), resulting in lower levels of HDL-C in the setting of high TG levels. The apo B-containing lipoproteins, now enriched in CE, are taken up by the liver receptors, as previously described. ⁷⁷ This exchange through CETP action is thought to be responsible for the inverse relationship between levels of TG and HDL-C. ⁷⁹ The cardioprotective effect of HDL has been largely attributed to its role in reverse cholesterol transport. HL then hydrolyzes the TGs within the TG-rich LDL to release free fatty acids, a process that remodels the LDL particles into smaller and denser LDL particles that can enter the arterial intima more easily than larger LDL particles, thus making them more atherogenic (see Fig. 15-2). Small, dense LDL particles also bind less avidly to the LDL receptor, thus prolonging their half-life in the circulation, making these particles more susceptible to oxidative modification and to subsequent uptake by the macrophage scavenger receptors. ⁸⁰

Using stable isotopes in the fed (nonfasting) state in humans, Welty and colleagues 81 showed that apo AI FCR is inversely correlated with the FCR of apo B48 (see Fig. 15-2 for details) but not with VLDL apo B100 FCR or production rate. Thus, when CM apo B48 clearance is delayed (represented by decreased apo B48 FCR in Fig. 15-2), TG-rich apo B particles accumulate and the TG is transferred to HDL apo AI particles in exchange for CE. These results suggest that in the fed state, levels of TG-rich apo B48 of intestinal origin are more important determinants of levels of HDL-C than the amount of TG-rich lipoproteins of hepatic origin, which contain apo B100.

Mutations in Enzymes of VLDL and Chylomicron Metabolism Affecting Risk for Atherosclerosis

Alterations in the enzymes involved in lipid metabolism may affect the risk for atherosclerosis. As noted earlier, LPL -hydrolyzes TGs contained in the core of CMs and VLDL. In partial LPL deficiency, TG-enriched lipoproteins have a prolonged circulation time that allows more interaction with the endothelium. An Asn291Ser substitution in LPL causes impaired function of this enzyme and is associated with an increase in plasma TG. Female but not male carriers of this mutation have a twofold increase in the risk of CHD and nonfatal cerebrovascular disease. 82,83

Apo C-II is an activator of LPL that hydrolyzes the core TGs, thereby releasing free fatty acids and making the CMs and VLDL progressively smaller and forming remnants. Apo C-II thus increases the catabolism of both CM and VLDL, thereby lowering TG levels. Apo C-III is an inhibitor of LPL, and in contrast to apo C-II, which lowers TG levels, apo C-III can raise TG levels by stimulating VLDL synthesis, inhibiting LPL, and inhibiting the binding of remnants to the LDL receptor mediated by apo E. Thus, high levels of apo C-III are associated with TGenriched VLDL particles that are ultimately converted to TG-rich remnants. These remnants are then lipolyzed to small, dense LDL particles by HL. 59 Thus, high levels of apo C-III are associated with TG-rich VLDL particles that circulate longer and are therefore ultimately converted to TG-remnants, which are then lipolyzed by HL to small, dense LDL particles. Most patients with elevated levels of VLDL have excess amounts of both TGs and apo C-III within their lipid particles. Persons lacking apo C-III have efficient lipolysis of TGs and therefore low levels of TG. 60

Produced in the liver, apo AV activates proteoglycan-bound LPL and thus accelerates TG hydrolysis from VLDL and CMs independently of other apoproteins. 84 A sequence element between residues 185 and 228 functions in binding of apo AV to heparin sulfate proteoglycans, members of the LDL receptor family and GPIHBP1. 85 Plasma levels of apo AV are extremely low, and this factor, plus the association of apo AV with cytosolic lipid droplets, suggests that apo AV may modulate TG metabolism within the cell. 85 The gene for apo AV is located at the apo A1/C3/A4/A5 gene cluster on chromosome 11q23. Several single-nucleotide polymorphisms are associated with significantly higher plasma TG levels in patients (ie, - 1131T > C, S19W, G185C). 84 The structural mutations Q139X, Q148X, and IVS3 + 3G > C predispose to familial hypertriglyceridemia and late-onset chylomicronemia. 84 Thus, apo AV is an important regulator of plasma TG levels in humans.

Polymorphisms in HL may also affect risk for atherosclerosis . There are four common sequence polymorphisms in the HL gene promoter; the most frequent is a C to T substitution . ⁸⁶ The presence of a C allele is associated with higher HL activity; smaller, denser, and more atherogenic LDL particles; and lower levels of HDL-C. ⁸⁷

CLASSIFICATION AND CAUSES OF HYPERTRIGLYCERIDEMIA

Hypertriglyceridemia can occur as an isolated hypertriglyceridemia or in combination with hypercholesterolemia (familial combined hyperlipoproteinemia and familial dysbetalipoproteinemia) (Table 15-1). Both forms can be further subdivided into primary and secondary causes. LPL, HL, and their apolipoproteins regulate levels of TG; therefore, abnormalities in any of these can affect TG levels.

Isolated Hypertriglyceridemia

Familial Hypertriglyceridemia—Severe

Familial hypertriglyceridemia is characterized by elevated TG levels with normal cholesterol and can be divided into severe and mild forms (see Table 15-1). In the severe form, TGs exceed 1000 mg/dL because of increases in both CMs and VLDL particles ^{25,88,89} (type V in the original Fredrickson-Levy classification, also called primary mixed hypertriglyceridemia). ²⁵ Most patients with mixed hypertriglyceridemia have familial hypertriglyceridemia due to partial deficiency of LPL or apo C-II (the ligand for LPL on CMs and VLDL) deficiency exacerbated by one or more of the secondary disorders noted in Table 15-1.

The other major primary cause of TGs > 1000 mg/dL is exogenous hyperlipemia or familial chylomicronemia due to CMs (hyperlipoproteinemia type I in the original Fredrickson-Levy classification). ²⁵ The most common primary cause of type I is complete absence of either LPL activity or apo C-II. 1,90,91 When LPL is absent (prevalence is 1 in a million), TG is generally > 2000 15 mg/dL. Patients with TGs > 2000 mg/dL usually have both a genetic form of hypertriglyceridemia and a secondary cause. Similar clinical manifestations of both types I and V include hepatosplenomegaly and occasional eruptive xanthomas (see Table 15-1). Features that distinguish type I from type V include (1) presentation of type I in childhood and of type V in adulthood; (2) absence of LPL or apo C-II activity or homozygous gene mutations in type I; (3) presence of a secondary factor in type V (alcohol, obesity, type 2 diabetes mellitus, hypothyroidism, or poor diet); (4) higher prevalence of type V than of type I; and (5) increased CMs alone in type I compared with elevations in both CMs and VLDL in type V.

The primary risk associated with TG levels > 1000 mg/dL is pancreatitis. Minimal atherosclerotic risk is reported for patients with hyperchylomicronemia (type I) or the severe form of familial hypertriglyceridemia (type V), probably because the lipoprotein particles are too large to enter the arterial wall. ²⁴

Patients with marked hypertriglyceridemia (> 1000 mg/dL [11.3 mmol/L]) may develop the chylomicronemia syndrome. This can include recent memory loss, abdominal pain or pancreatitis, dyspnea, eruptive xanthoma, flushing after alcohol ingestion, and lipemia retinalis. 90

Familial Hypertriglyceridemia—Mild

The mild form of familial hypertriglyceridemia (type IV hyperlipoproteinemia phenotype) is an autosomal dominant disorder characterized by mild to moderate elevations in TG from 200 to 500 mg/dL, often in association with insulin resistance, obesity, hyperglycemia, hypertension, hyperuricemia, and low HDL-C. Mutations in the LPL gene decrease enzyme activity and therefore delay the degradation of CMs and VLDL that carry endogenous TGs. Gly188Glu, Asp9Asn, and Asn291Ser have N-terminal mutations that reduce the activity of LPL, resulting in an increase in serum TGs by 20% to 80% and also lower levels of HDL-C. ^{1.91} More marked elevations require some other factor, such as one of the drugs or acquired disorders (eg, estrogen replacement therapy in postmenopausal women).

Familial hypertriglyceridemia is common in patients with premature CHD. The prevalence of familial hypertriglyceridemia with low HDL-C levels was 15% in patients undergoing coronary arteriography before the age of 55 years. 93 Among first-degree relatives of affected patients, baseline serum TG levels predicted cardiovascular mortality, independent of serum total cholesterol. 94



Classes

Type IV

5%-10%

Type V

1 X 10 3

Presents in childhood

pancreatitis,

or adolescence;

abdominal pain,

hepatosplenomegaly,

(T chylomicrons with

i I_DL and f HDL)

Endogenous hyperlipemia

hypertriglyceridemia

hyperlipemia (T VLDL +

T chylomicrons)

(T VLDL)

Primary mixed

Family combined

and HDL)

hyperlipoproteinemia

(TVLDL and TLDL and

Family

TG > 1000 mg/dL Familial lipoprotein lipase deficiency or familial apo C-II deficiency (95% of cases); LPL

Lipid Levels

TG: 200-999 mg/dL

TG > 1000 mg/dL

mutations 105,108; apo C-II mutations 105 (rare)

Family multiple

lipoprotein type hyperlipidemia

Dysglobulinemias 5 Family hypertriglyceridemia Systemic lupus (mild form)

Diabetic hyperlipemia ...s Glycogenosis, type I s Lipodystrophy Uremia

(Estrogen use)

(Glucocorticoid use) 5

(Stress-induced) 5

Dysglobulinemias

ervthematosus

Systemic lupus

Sporadic hypertrialyceridemia Tangier disease Hypopituitarism No replicated causative or Nephrotic syndrome 5 (Diabetes mellitus) +, 5 susceptibility genes Family (Alcoholism) =

(severe form) Familial lipoprotein lipase deficiency; heterozygous LPL mutations (5%-10% of cases) or apo C-II deficiency

hypertrialyceridemia

retinal lipemia Asymptomatic; may be associated with increased risk of erythematosus : vascular disease

> Present in adulthood: hepatosplenomegaly, eruptive xanthomas, abdominal pain,

pancreatitis: lipemia retinal

Tuberous gold

Hypertriglyceridemia and Hypercholesterolemia

Remnant hyperlipidemia Type III IG variable: can be (O0 -VLDL) 1 X 10 2: expression is 1 x 10 4

Type IIb

> 1000 mg/dLTC = 250-500 mg/dL

TG: 200-500 mg/dL

TC: 200-400 ma/dL:

small, dense LDL

dvsbetalipoproteinemia Homozygous APOE2 mutation plus secondary genetic or medical factors

Obligate heterozygotes for mutations of LPL or apo C-III; insulin resistance Autosomal dominant USF1 105

Hypothyroidism Systemic lupus ervthematosus

Nephrotic syndrome

Cushing's syndrome

(Glucocorticoid use) (Stress-induced)

Hypothyroidism

Dysglobulinemias

tuberoeruptive xanthomas on extensor surfaces of extremities; plantar palmar crease xanthomas; vascular disease

Vascular prematurity disease

Hypertriglyceridemia and Hypercholesterolemia

Hypertriglyceridemia can occur in two phenotypes in combination with hypercholesterolemia. The first is familial combined hyperlipoproteinemia (FCHL), and the second is familial dysbetalipoproteinemia.

Familial Combined Hyperlipoproteinemia

In FCHL (type IIb), overproduction of hepatically derived VLDL apo B100-containing lipoproteins results in plasma TG levels of 200 to 500 mg/dL and plasma cholesterol levels of 200 to 400 mg/dL and small, dense LDL. 95,96 FCHL has an autosomal dominant mode of inheritance with variable pen etrance and a population prevalence of 1% to 2%. 97 It is the most common familial lipid disorder in post-MI patients 98 and accounts for one third to one half of familial causes of CHD. 94 The molecular basis includes mutations in LPL and apo C-III and upstream stimulatory factor 1 (USF1), which encodes an upstream stimulatory factor. 99,100 FCHL is also linked with insulin resistance 101,102 due to both increased free fatty acid flux from the periphery and insulin-stimulated lipogenesis, which increases the production of VLDL. 103 Families with premature CHD often have either familial dyslip idemia (high TGs and low HDL-C [type IV]) or FCHL (high TGs, high LDL-C, and low HDL-C [type

IIb]). 104

Patients with FCHL can present with combined hypercholesterolemia and hypertriglyceridemia or either abnormality alone. Thus, subjects with FCHL who overproduce VLDL particles and also synthesize TG at an increased rate will secrete an increased number of large, TG-rich VLDL particles. If they are unable to efficiently catabolize these VLDL particles because of an LPL mutation or low LPL activity, they will have a high TG level but a normal or reduced number of LDL



particles and thus a normal LDL-C level. On the other hand, with efficient catabolism of the increased numbers of large, TG-rich VLDL particles, the number of LDL particles is increased, resulting in both increased TG and LDL-C levels. Finally, subjects who synthesize a normal amount of TG but an increased number of VLDL particles (with normal TG load) have increased numbers of LDL particles and elevated plasma LDL-C levels but a normal TG level. LPL may be responsible for part of this phenotypic variability as hypertriglyceridemia is more prominent in patients with LPL deficiency ¹⁰⁵ or an LPL gene mutation. ¹⁰⁶

Familial Dysbetalipoproteinemia

As noted before, the apo E ligand is necessary for receptormediated catabolism of CM and VLDL remnants. There are three isoforms of the apo E allele: apo E2, apo E3, and apo E4. Apo E3/3 is the most common apo E genotype. Apo E2 differs from apo E3 by substitution of cysteine for the normal arginine at residue 158 in the receptor-binding domain. Consequently, apo E2 does not bind as well as apo E3 to the apo B/E (LDL receptor). ¹⁰⁷ Subjects with familial dysbetalipoproteinemia have the apo E2/E2 genotype, which is an autosomal recessive disorder characterized by an accumulation of VLDL and CM remnants in the plasma due to inefficient uptake through apo E2, which binds poorly to hepatic LDL-related receptors. ¹⁰⁸ Consequently, they have an increase in cholesterol-enriched VLDL (\$ -VLDL, also termed IDL) and CM remnants. The prevalence of the apo E2 isoform is 1 in 100; however, only approximately 1 in 10,000 carriers exhibits the dyslipidemia, which is thought to be triggered by a secondary cause, such as marked hyperglycemia (type 2 diabetes), hyperuricemia (gout), hypothyroidism, or obesity. 108 Classic physical findings include tuberoeruptive xanthomas and xanthomas of the palmar creases; the risk for CHD is increased.

Secondary Causes

The major secondary causes (see Table 15-1) of hypertriglyceridemia include poorly controlled type 2 diabetes, obesity, excessive alcohol intake, renal disease, pregnancy, medica tions (Table 15-2), excessive ingestion of saturated fats and simple sugars, nonalcoholic hepatosteatosis, and physical inactivity. Alcohol intake can cause elevated TG by inhibition of LPL ¹⁰⁹ or increased VLDL TG production. For every gram of alcohol consumed per day, TG concentration can increase on average by 0.19 mg/dL, which is about 5.7 mg/dL for 30 g of alcohol. ¹¹⁰ These secondary causes must always be considered during the evaluation and treatment of hypertriglyceridemia.

Metabolic Syndrome

The association of hypertriglyceridemia with obesity and diabetes or glucose intolerance is termed the metabolic syn drome, which is defined clinically according to the National Cholesterol Education Program 111 by at least three of the following: central obesity (waist circumference > 35 inches in women and > 40 inches in men), fasting blood glucose concentration > 100 mg/dL, TGs > 150 mg/dL, low HDL-C (<40 mg/dL in men and < 50 mg/dL in women), and systolic or diastolic blood pressure > 130/ > 85 mm Hg. Atherogenic dyslipidemia in metabolic syndrome and people with type 2 diabetes (termed diabetic dyslipidemia) is characterized by elevated TGs and small, dense, cholesterol-depleted LDL and HDL particles. 111 Insulin resistance increases mobilization of free fatty acids from adipose tissue to liver, where increased production of VLDL occurs; thus, hypertriglyceridemia in type 2 diabetes and metabolic syndrome is usually secondary to increased VLDL concentrations in plasma, with or without chylomicronemia. 112 Downregulation of LPL expression in

insulin resistance leads to decreased catabolism of TG-rich

TABLE 15-2	Effects of Selected Drugs on Triglyceride and Cholesterol Levels*			
Drug		Triglycerides	LDL-C	HDL-C
Alcohol		Increased	No effect	Increased
Estrogens, estrad	iol	Increased	Decreased	Increased
Androgens, testosterone		Increased	Increased	Decreased
Progestins		Decrease	Increase	Decrease
Glucocorticoids		Increased	No effect	Increased
Cyclosporines		Increased	Increased	Increased
Tacrolimus		Increased	Increased	Increased
Thiazide diuretics		Increased	Increased	Decreased
Beta blockers		Increased	No effect	Decreased
Sertraline		Possible increase	Increased	No effect
Protease inhibito	rs	Increased	No effect	No effect
Valproate and related drugs		Increased	No effect	Decreased
Isotretinoin		Increased	No effect	Decreased
Clozapine, olanzapine [,]		Increased	No effect	Decreased

*Alcohol, estrogens, estradiol, glucocorticoids, thiazide diuretics, beta blockers, sertraline, protease inhibitors, valproate and related drugs, and isotretinoin can cause severe hypertriglyceridemia and the chylomicronemia syndrome in patients with a familial form of hypertriglyceridemia.

Second-generation antipsychotics: clozapine and olanzapine have most effect; risperidone and quetiapine have intermediate effects; and aripiprazole and ziprasidone have least effect.

Data from Brunzell JD: Clinical practice. Hypertriglyceridemia. *N Engl J Med* 357:1009, 2007. Copvright © 2007 Massachusetts Medical Society. All rights reserved. VLDL. The higher levels of TG promote CETP-mediated transfer of CE from HDL, thus producing TG-rich small, dense HDL that are catabolized more rapidly, leading to low levels of HDL-C. Small, dense HDL also have reduced antioxidant and anti-inflammatory properties. The metabolic syndrome and increased TG-rich lipoproteins are also associated with a pro-inflammatory and prothrombotic state due to the presence of atherogenic lipoproteins, clotting factors, and increased plasma viscosity. ¹¹³ In clinical practice, elevated serum TGs are most often observed in persons with the metabolic syndrome, although secondary or genetic factors can raise TG levels. Metabolic syndrome has a prevalence of 24% in US adults and 43% in adults older than 60 years ¹¹⁴; therefore, it is a major health problem.

LABORATORY EVALUATION OF HYPERTRIGLYCERIDEMIA

In this section, the literature on several clinical laboratory approaches to assess the risk of TGs and TG-rich lipoproteins is reviewed.

Fasting Versus Nonfasting Triglyceride Levels

The Friedewald equation is often used to estimate LDL-C levels: LDL-C = TC - (TG/5 + HDL-C); it requires

238 measurement of fasting TG levels. TG/5 is an estimate of the cholesterol in VLDL and IDL particles (VLDL-C + IDL-C). This equation has been used for more than 40 years to calculate LDL-C; therefore, the majority of research studies have examined the association of TG with CHD on the basis of fasting TG levels. As shown earlier in the section on lipoprotein metabolism, levels of TG-rich- and apo B48containing CMs of intestinal origin are probably more important determinants of levels of HDL-C than the amount of TG-rich lipoproteins of hepatic origin, which contain apo B100. 81 In the fasting state, very little apo B48 is being produced; therefore, it would make sense that nonfasting levels of TG might be a better predictor not only of the concentration of TG-rich lipo proteins circulating in plasma most of the time but also of the HDL-C level.

In 2007, Bansal and colleagues 115 and Nordestgaard and coworkers 116 reported that in two long-term prospective cohort studies, TG levels obtained 2 to 4 hours postprandially predicted risk of CHD better than TG levels measured after a 12- to 14-hour fast and better than LDL-C calculated by the Friedewald equation. 115-117 In both studies, elevated postprandial TG levels increased the risk of CHD for both sexes; however, women had a greater risk of CHD associated Much data supports apo B as a predictor of CHD. 131,132 Fred rickson postprandial increases in TG levels.

fresh for each test. Second, a fat load may cause nausea or laboratory assays in hospital chemistry laboratories. vomiting. Third, requirement for a 2- to 4-hour postpran dial peak may not be practical in outpatient care. In the long run, Non-HDL-C fasting TG measurements may be more reliable because of controlled conditions. The strong correlation between postprandial and fasting TG levels may obviate the need for repeated postprandial measurements. 119

Lipoprotein Particle Subclasses

correlation with TGs. In the Veterans Affairs High-Density achieved (Table 15-3). 111 Lipoprotein Intervention Trial (VA-HIT), a change in TG concentration did not predict the magnitude of reduction in PREDICTIVE VALUE OF NON-HIGHrisk of CHD events; rather, it was the change in LDL and HDL particles that predicted change in risk. 125

Patients with CHD often have increases in small, dense HEART DISEASE LDL; however, LDL particle size is not an independent predictor of CHD, and in fact, the major factor regulating LDL Several observational and intervention studies have reported that particle size is plasma TG level. 126 Therefore, the measurement elevated levels of non-HDL-C are predictive of of particle size has not been proven to provide better information than standard lipid and lipoprotein measurements. Patients with TG levels > 150 mg/dL generally have increased levels of small, dense LDL particles. 127 As

noted earlier, after conversion of large VLDL particles to TGrich remnants, HL hydrolyzes the remnants to small, dense LDL particles. 128 Consequently, levels of TG and VLDL are strongly and positively correlated with levels of small, dense LDL particles. ¹²⁷ In some prospective, nested case-control studies, subjects with small, dense LDL particles have an increased risk for CHD; however, other studies have concluded that increased levels of small LDL particles are not an independent predictor of CHD but rather are products of elevated levels of TG-rich lipoproteins that are associated with an atherogenic milieu-elevated TG, reduced HDL-C, and potentially other biomarkers of the metabolic syndrome. 129 Because both large and small LDL subtypes have been shown to predict CHD, apo B concentrations, which estimate particle numbers, rather than LDL particle size, may be the better predictor of CHD. 120 Moreover, apo C-III enrichment of apo Bcontaining lipoproteins has been linked to the atherogenicity of apo B-containing lipoproteins and therefore may also be a better predictor of CHD than particle size. 130

Apo B

with hypertriglyc eridemia than men did, 115,116 confirming and associates 133 recognized more than 40 years ago that prior studies that showed a higher risk in women than in men. atherosclerosis is more closely related to the total number of apo B-14,118 Postprandial lipoproteins are TG rich, and if their containing particles rather than to LDL-C or TG concentrations catabolism is delayed (insulin resistance, LPL mutations), the alone. One apo B molecule is present on the surface of VLDL, IDL, products of their metabolism, CM remnants and small dense LDL, and lipoprotein(a), a molecule of apo B100 covalently bound to LDL, can remain in the plasma for 12 hours or more, 24 with apoprotein (a) 25; therefore, the apo B level may provide a more exposure of the endothelium to TG-rich, atherogenic remnant direct measure of circulating atherogenic lipoproteins. 134 As noted particles, a finding accounting for the greater CHD risk with earlier, not all forms of hypertriglyceridemia are atherogenic 24; however, the relative atherogenicity of apo B is well established by Although postprandial TG levels may predict risk better the fact that modification of lysine residues on apo B is necessary for than fasting TG levels, nonfasting TG measurements may be uptake by scavenger receptors on macrophages. 52-54 However, the difficult to incorporate into clinical practice. First, a standard routine measurement of apo B is not always practical because of cost fat-feeding protocol would need to be developed and prepared and technical limitations that preclude me surement in routine

For all of these reasons, as suggested by the National Choles terol Education Program (NCEP) Adult Treatment Panel III (ATP III), 111 non-HDL-C is a practical and useful surrogate measure of atherogenic particle concentration 120,135 and more predictive of CHD risk than LDL-C level alone (especially when TG levels are elevated) because this measure is a sum of all atherogenic lipoproteins. 106,111 Lipoprotein particle composition has been associated with Non-HDL-C is TC - HDL-C, which is the sum of VLDL-C, IDL-C, differences in the relative atherogenicity of lipoproteins. 120,121 and LDL-C. Therefore, non-HDL-C includes the cholesterol in all of Small, dense LDL particles are more susceptible to accelerated the atherogenic apo B-containing lipoproteins: TG-enriched oxidation and are incorporated more readily by vascular wall lipoproteins, CMs, CM remnants, VLDL and VLDL remnants, IDL, macrophages than other lipoprotein particles are. 81,122 Several LDL, and lipoprotein(a). Non-HDL-C is accurate and reliable in a prospective cohort studies have reported that the number of nonfasting state; therefore, measurement of non-HDL is practical small, dense LDL particles is a greater predictor of CHD risk and easy. 111 When lipid levels are normal, non-HDL-C is highly than are measured levels of serum LDL-C. 123,124 Thus, unlike correlated with apo B levels. 136 Because high LDL-C and TG levels the linear relationship of risk with LDL-C, the risk of CHD confer greater risk for CHD than high LDL-C alone, NCEP ATP III associated with elevated TG levels may be a function of the guidelines recommend non-HDL-C as a secondary target of therapy associated lipoprotein disorder more than a direct numerical when the serum TG level is > 200 mg/dL after the LDL-C target is

LIPOPROTEIN CHOLESTEROL FOR CORONARY

Risk Category	LDL-C Goal	Non-HDL-C Goal +
Very high risk _§ ··	< 70 mg/dL (optional)	< 100 mg/dL
High risk: CHD § or CHD risk equivalents "	< 100 mg/dL	< 130 mg/dL
Moderately high risk: > 2 risk factors ^ (10-year risk 10%-20%)	< 130 mg/dL	< 160 mg/dL
Moderate risk: > 2 risk factors (10-year risk < 10%)	< 130 mg/dL	< 160 mg/dL
Lower risk: 0 or 1 risk	< 160 mg/dL	< 190 mg/dL

*NCEP ATP III guidelines for non-HDL-C state that in addition to the primary goal of LDL-C reduction, non-HDL-C is a secondary target of therapy in patients with TG levels of 200 to 499 mg/dL. Because a normal VLDL-C level is <30 mg/dL, the therapeutic goal for non-HDL-C is 30 mg/dL higher than the goal for LDL-C. Factors that place patients at very high risk in favor of a decision to reduce LDL-C levels to <70 mg/dL. The optional goal of <70 mg/dL does not apply to subjects who are not very high risk.

When TGs are 200 to 499 mg/dL.

- ^ Presence of established cardiovascular disease plus (1) multiple major risk factors (especially diabetes mellitus), (2) severe and poorly controlled risk factors (especially continued cigarette smoking), (3) multiple risk factors for the metabolic syndrome (especially high TGs [>200 mg/dL] plus non-HDL-C >130 mg/dL with low HDL-C [<40 mg/dL]), and (4) acute coronary syndromes.
- § CHD includes history of myocardial infarction, unstable angina, stable angina, coronary artery procedures, or evidence of clinically significant myocardial ischemia.
- || CHD risk equivalents include clinical manifestations of noncoronary forms of atherosclerotic disease, diabetes, and >2 risk factors, with 10-year risk for hard CHD >20%
- ^Risk factors include cigarette smoking, hypertension (blood pressure <140/90 mm Hg or taking antihypertensive medication), low HDL-C (<40 mg/dL), family history of premature CHD, and age (men <45 years, women <55 years).

From Miller M, Ginsberg HN, Schaefer E: Relative atherogenicity and predictive value of nonhigh-density lipoprotein cholesterol for coronary heart disease. Am J Cardiol 101:1004. 2008, with permission from Elsevier Limited.

and women with LDL-C levels < i00 mg/dL in the Atherosclerosis nonfasting state. Risk in Communities (ARIC) study; elevated TG levels were associated with substantially greater relative risks in women (4.7) than in men (2.i) after adjustment for LDL-C, HDL-C, and lipoprotein(a)? 40 In prospective follow-up of a cohort of i5,632

ATP III presented the following classification of serum TG levels" 4:

- Normal: < i50 mg/dL (i.7 mmol/L)
- Borderline-high: i50-i99 mg/dL (i.7 to 2.2 mmol/L)
- · High: 200-499 mg/dL (2.2 to 5.6 mmol/L)
- \sim Very high: > 500 mg/dL (> 5.6 mmol/L)

healthy women older than 45 years baseline in the Women's Health Study, non-HDL-C was as good as or better than apolipoprotein fractions for prediction of risk of a first cardiovascular event?" In 6year follow-up of i8,225 men in the Health Professionals Follow-up Study free of CHD at baseline, the relative risk of CHD in the highest quintile of non-HDL-C compared with the lowest quintile was 2.76 after adjustment (95% CI, i.66-4.58), which was better than LDL-C, i.8i (95% CI, i.i2-2.93), but not quite as good as apo B, 3.0i (95% CI, i.8i-5.00). After mutual adjustment of non-HDL-C and LDL-C, only non-HDL-C was predictive of CHD? 37 In prospective follow-up of i562 men and i760 women older than 30 years and free of CHD at baseline in the Framingham Heart Study, the predictive value of non-HDL-C was better than LDL-C and comparable to apo B for prediction of risk of CHD. i32 After multivariate adjustment in a subsequent analysis of the combined original Framingham Heart Study cohort and the offspring (2693 men, 3i0i women), VLDL-C was an independent predictor of risk and non-HDL-C level was a stronger predictor of CHD risk than LDL-C alone at TG levels both greater than and less than 200 mg/dL? 42 These results suggest that VLDL-C contributes to the development of CHD in addition to LDL-C and thus support the fact that both VLDL-C and LDL-C are essential in predicting CHD risk. In fact, non-HDL-C was better than

The Emerging Risk Factors Collaboration analyzed records of 302,430 people without initial vascular disease from 68 long-term prospective studies (Europe and North America) for a total of i2,785 cases of CHD (8857 nonfatal MIs and 3928 deaths due to CHD) during 2.79 million person-years of follow -up (median, 6.i years to first outcome)? 43 The hazard ratio for the primary outcome (nonfatal MI and CHD death) for TG was i.37 (95% CI, i.3i-i.42) after adjustment for non lipid risk factors. However, after further adjustment for HDL-C and non-HDL-C, the hazard ratio for TG was reduced to 0.99 (95% CI, 0.94-i.05) (Fig. i5-3). The hazard ratio for CHD with non-HDL-C was i.56 (95% CI, i.47-i.66) after adjustment for nonlipid risk factors. After adjustment for HDL-C and log TG, the hazard ratio for non-HDL-C remained significant at i.50 (95% CI, i.39-i.6i) (Fig. i5-4). There was no difference between those who fasted and those who were nonfasting and no difference by gender for either TG or non-HDL-C. In a subset with available measurements, the hazard ratio for directly measured LDL-C was i.38 (95% CI, i.09-i.73), which was similar to the hazard ratio of i.42 cardiovascular disease and cardiovascular disease mortality, 131,132,137,138 (95% CI, i.06-i.9i) for non-HDL-C in the same subset. Analysis by similar to the predictive value of apo B and as good as or better than `HDL-C showed that after adjustment for nonlipid risk factors, nonthat of LDL-C. In the Bypass Angioplasty Revascularization HDL-C, and log TG, the hazard ratio for CHD with HDL-C was 0.78 Investigation (BARI), 1514 patients with multivessel CHD were (95% CI, 0.74-0.82). When analyzed by quintiles of HDL-C levels, the observed for 5 years. 13 9 Non-HDL-C, but not HDL-C or LDL-C, was hazard ratio was 0.35 (95% CI, 0.30-0.42) for a 15 mg/dL higher HDLa significant univariate and multivariate predictor of nonfatal MI C and 80 mg/dL lower non-HDL-C; this was not changed by (RR, i.049; 95% CI, i.006-i.093; P< 0.05) and angina pectoris (RR,i.049; addition of TĞ level. The hazard ratios for non-HDL-C and apo B 95% CI, i.004-i.096; P < 0.05) at 5 years. 13 9 Non-HDL-C did not predict were very similar in magnitude and shape, as were those for HDL-C mortality in this study. In long-term follow-up from the Lipid and apo AI (Fig. i5-5), findings suggesting that apolipoprotein Research Clinics Program Follow-up Study, increases of 30 mg/dL measurements are no better in predicting risk than cholesterol levels. in non-HDL-C and LDL-C levels cor responded to increases in CVD i43 In summary, this largest meta-analysis suggests that TG level mortality of i9% and i5%, respectively, in men and ii% and 8%, provides no additional information about risk when non-HDL-C is respectively, in women?" The risk for CHD death was lowest in men calculated from TC and HDL-C levels in either the fasting or





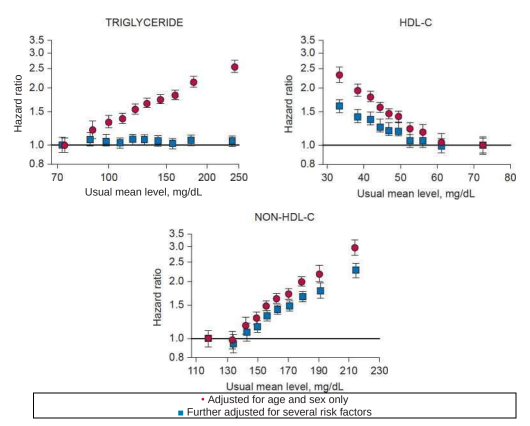
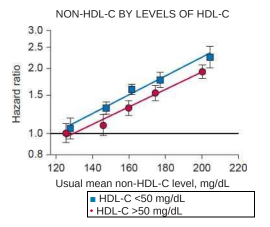


FIGURE 15-3 Hazard ratios for coronary heart disease across quantiles of usual triglyceride, HDL-C, and non-HDL-C levels. Analyzes for coronary heart disease were based on 302,430 participants (involving 12,785 cases) from 68 studies. Regression analyzes were stratified, where appropriate, by sex and trial group. Values with further adjustments were adjusted for age, systolic blood pressure, smoking status, history of diabetes mellitus, and body mass index; furthermore, analyzes of log otriglyceride were adjusted for HDL-C and non-HDL-C levels, analyzes of HDL-C were adjusted for non-HDL-C and log otriglyceride levels, and analyzes of non-HDL-C were adjusted for HDL-C and log otriglyceride levels. Studies with fewer than 10 cases were excluded from analysis. Sizes of data markers are proportional to the inverse of the variance of the hazard ratios. The *y*-axes are shown on a log scale. The *x*-axes for triglycerides are shown on a log scale. Referent groups have lowest quantiles for triglyceride and non-HDL-C and highest quantiles for HDL-C. Error bars indicate 95% confidence intervals. (From Di Angelantonio E, Sarwar N, Perry P et al; Emerging Risk Factors Collaboration: Major lipids, apolipoproteins, and risk of vascular disease. JAMA 302:1993, 2009. Copyright © 2009 American Medical Association. All rights reserved.)



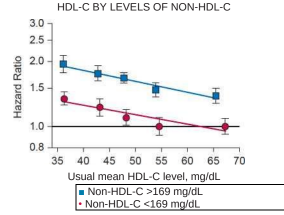
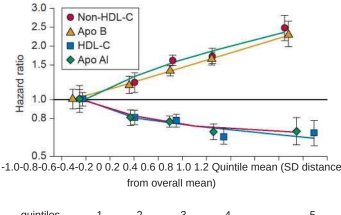


FIGURE 15-4 Hazard ratios for coronary heart disease across fifths of non-HDL-C by levels of HDL-C and fifths of HDL-C by levels of non-HDL-C. Analyses were based on 302,430 participants (involving 12,785 cases) from 68 studies, Median values in the Emerging Risk Factors Collaboration were 50 mg/dL for HDL-C and 169 mg/dL for non-HDL-C. Regression analyses were stratified, where appropriate, by sex and trial group and adjusted for age, systolic blood pressure, smoking status, history of diabetes mellitus, body mass index, and loge triglyceride levels. Studies with fewer than 10 cases were excluded from analysis. Sizes of data markers are proportional to the inverse of the variance of the hazard ratios. The *y*-axes are shown on a log scale. Referent groups are lowest fifth of non-HDL-C in the higher level of HDL-C and highest fifth of HDL-C in the lower level of non-HDL-C. Lines are fitted by log-linear regression of log hazard ratios on mean levels. Error bars indicate 95% confidence intervals. (From Di Angelantonio E, Sarwar N, Perry P et al; Emerging Risk Factors Collaboration: Major lipids, apolipoproteins, and risk of vascular disease. JAMA 302:1993, 2009. Copyright ⊚2009 American Medical Association. All rights reserved.)



quintiles	1	2	3	4	5
Mean usual level, mg/dL					
Non-HDL-C	125	145	159	173	198
Аро В	85	99	108	118	137
HDL-C	37	44	49	55	66
About AL	126	139	148	158	178

FIGURE 15-5 Hazard ratios for coronary heart disease across fifths of usual lipids or apolipoproteins. Analyzes were based on 91,307 participants (involving 4499 cases) from 22 studies. Regression analyzes were stratified, where appropriate, by sex and trial group and adjusted for age, systolic blood pressure, smoking status, history of diabetes mellitus, and body mass index; furthermore, analyzes of non-HDL-C were adjusted for HDL-C and log e triglyceride, analyzes of apo B were adjusted for apo Al and log etriglyceride, analyzes of HDL-C were adjusted for non-HDL-C and log a triglyceride, and analyzes of apo AI were adjusted for apo B and log a triglyceride. Studies with fewer than 10 cases were excluded from analysis. Sizes of data markers are proportional to the inverse of the variance of the hazard ratios. Referent groups have the lowest fifths. Lines are fitted by first-degree fractional polynomial regression of log hazard ratios on mean SD score. Error bars indicate 95% confidence intervals. The y -axis is shown on a log scale. The X -axis is shown on a Z-transformed scale. (From Di Angelantonio E, Sarwar N, Perry P, et al; Emerging Risk Factors Collaboration: Major lipids, apolipoproteins, and risk of vascular disease. JAMA 302:1993, 2009. Copyright © 2009 American Medical Association. All rights reserved.)

have shown that CHD risk begins to increase at a fasting TG concentration of 160 to 190 mg/dL (1.8 to 2.2 mmol/L), a finding that provides support for a VLDL-C level < 30 mg/dL being normal (or a TG level < 150 mg/dL being normal). 16,22,144 Moreover, the association of significant reduction in risk with a TG level < 150 mg/dL in the PROVE-IT trial (discussed earlier) provides further support that a VLDL-C level < 30 mg/dL is desirable. 23 The results of the COLTS study discussed earlier suggest that risk increases in patients with CHD at TG levels above 100 mg/dL (1.13 mmol/L), 15 a finding supporting the use of the non-HDL-C goal of < 100 mg/dL when the optional goal of LDL-C < 70 mg/dL is used in patients with CHD.

Table 15-4 outlines a management approach to patients with hypertriglyceridemia. In evaluation of a patient with hypertriglyceridemia, thyroid-stimulating hormone, fasting glucose concentration, blood urea nitrogen, and creatinine should be measured to exclude potential secondary causes (see Table 15-1), and medications that may be raising TGs (see Table 15-2) should be stopped or switched if possible. According to the NCEP, the primary goal of therapy with borderline-high or high TG levels (150 to 499 mg/dL) is to achieve the LDL-C goal (see Table 15-3). 111 In primary prevention, lifestyle changes can be followed for at least 3 to 6 months. In secondary prevention, lifestyle changes and statins can be prescribed simultaneously. LDL-C goal is achieved, the following the recommendations should be observed for various levels of elevated TGs (see Table 15-4 for summary):

· When TG levels are borderline high (150 to 199 mg/dL [1.7 to 2.2 mmol/L]), emphasis is on therapeutic lifestyle changes (diet, weight reduction, and increased physical activity) as recommended by the NCEP. 111 When TG levels are high (200 to 499 mg/dL [2.2 to 5.6 mmol/L]), non-HDL-C becomes a secondary target of therapy after the LDL-C goal is achieved. If lifestyle changes do not reach the non-HDL-C goal in primary prevention, drug therapy should be considered in high-risk patients. In a consensus statement, statins remained the first-line therapy for moderate hypertriglyceridemia; the most effective TG-lowering drugs,

TABLE 15—4 Management Approach to Hypertriglyceridemia

TG 150-199 mg/dL

Counseling on lifestyle changes, diet, 30 minutes of daily aerobic exercise, and ideal body weight

TG 200-499 mg/dL; Non-HDL-C is Secondary Target After LDL-C Goal is Reached (See Table 15-3)

Review medications (see Table 15-2)

Change to lipid-neutral or favorable agents when possible (eg, alpha blockers, biguanides, thiazolidinedione).

Lower doses of or stop drugs that increase triglycerides, such as beta blockers (particularly nonselective agents), glucocorticoids, diuretics (thiazide and loop), ticlopidine, estrogens when indicated clinically.

Laboratory studies: exclude secondary disorders of lipid metabolism.

Fasting blood glucose concentration

Serum blood urea nitrogen and creatinine levels

Thyroid function studies (thyroid-stimulating hormone)

Therapeutic lifestyle changes (diet and exercise)

Weight loss

Avoid concentrated sugars and simple carbohydrates

Reduce saturated fat

Reduce or eliminate alcohol

Increased omega-3 fatty acid intake through fish consumption

Aerobic exercise minimum of 3 hours weekly

Recheck lipid profile in 3 to 6 months (give enough time for adequate weight loss).

Primary Prevention

If LDL-C goal is not reached in 3 to 6 months with steps 1 to 3, consider adding statin

Once LDL-C goal is reached, determine non-HDL-C goal,

Reinforces lifestyle changes. If still not at non-HDL-C goal, consider Niacin (extended release), especially if HDL-C is low; up to 2 g daily Fish oil (omega-3 fatty acids) up to 3.2 g EPA and DHA daily (especially if HDL-C is normal)

Repeat laboratory studies 6 to 8 weeks after dose adjustments.

Secondary Prevention

In addition to the following steps 1 to 3, statin therapy should be used to reach the LDL-C goal, with repeated laboratory studies 6 to 8 weeks after dose adjustments.

TG 500-999 mg/dL

Weight loss; increased exercise—follow steps 1 to 3.

I consider a very-low-fat diet (< 15% of calorie intake).

Remember that LDL-C cannot be estimated when TG > 400 mg/dL.

Consider fibrate therapy or extended-release niacin (eg, monitor INR on fibrate and

Fenofibrate

Fenofibrate micronized 67, 134, or 200 mg/day taken with dinner

Fenofibrate 54 or 160 mg/day; unnecessary to take with meal

Tricor (nanocrystallized) 48 or 145 mg/day; taken without regard to meals Gemfibrozil (Lopid) 600-1200 mg/day (usually 600 mg bid), 30 to 60 minutes before meals

Extended-release niacin

Niaspan, up to 2 g daily at bedtime: take with snack; aspirin 30 minutes before

TG > 1000 mg/dL

Follow steps 1 to 3.

Initiate fibrate therapy—monitor serum creatinine.

With acute pancreatitis:

Very-low-fat diet (10%-15% of energy intake) Cessation of alcohol

Insulin, if indicated for glycemic control

Admit patient to hospital if necessary.

Nothing by mouth; IV fluid replacement

Plasma exchange has been used.

fibrates and niacin, both of which also raise HDL-C (Table 15-5), were suggested for high-risk patients. 145 If patients in this category have low levels of HDL-C, nicotinic acid should be considered for its HDL-C-raising properties in addition to lowering of TG levels. When HDL-C level is normal, fish oil is another option to lower TG levels. Such patients may have a low or normal LDL-C level that increases as the serum TG concentration is reduced and may require the addition of or increase in a statin drug. 146

• When TG levels are very high (> 500 mg/dL [> 5.6 mmol/L]), the initial goal is to prevent pancreatitis by lowering TG levels with the combination of very-low-fat diets (< 15% of calorie intake), weight reduction, increased physical activity, and a TG-lowering drug such as a fibrate or nicotinic acid. Once TG levels are below 500 mg/dL, the LDL-C goals should be addressed. Although pancreatitis is infrequent until the TG level is > 1000 mg/dL, efforts to reduce the TG level to < 500 mg/dLmg/dL are important. 147

Adult patients with a TG level > 1000 mg/dL usually have a type V hypertriglyceridemia, although it can also result from secondary causes, such as diabetes, alcohol, and steroid hormones. Fasting glucose and thyroid-stimulating hormone concentrations must always be checked. These patients are at high risk for pancreatitis and need immediate treatment with both lifestyle changes and a fibric acid derivative. 148 Combination therapy with nicotinic acid may be necessary.

Diabetics. The American Diabetes Association has recognized the serum TG level as a surrogate for atherogenic TG-rich lipoproteins and recommends a target of < 150 mg/dL in the setting of baseline statin therapy, which is recommended for all diabetics. 149 Based on NCEP, the LDL-C goal for diabetics is < 100 mg/dL; therefore, the non-HDL-C goal would be < 130 mg/dL (or VLDL-C < 30 mg/dL, which is equivalent to a TG level of < 150 mg/dL). An optional LDL-C goal for diabetics with CAD can be < 70 mg/dL (see Table 15-3).

Treatment: Lifestyle Changes

Lifestyle changes-diet, exercise, and weight loss-are the cornerstone for treatment of elevated TG levels. In fact, medications can be ineffective in lowering TG levels if the patient does not make lifestyle changes, especially in patients with insulin resistance, in whom weight loss is essential to improve insulin resistance.

Dietary changes that can improve high TG levels (and low HDL-C levels in atherogenic mixed dyslipidemia of metabolic syndrome and type 2 diabetes) include a diet low in glycemic index (eliminating white bread, white potatoes, white rice, and soda) 150 and high in omega-3 fatty acids and fiber (vegetables, fruits, and beans). 151-156 NCEP and the American Heart Association recommend total fat intake ranging from 25% to 35% with < 7% saturated fat. 111,157 Polyunsaturated fat can range up to 10% in recognition of the beneficial effects of omega-3 fatty acids found in fatty fish and canola and soybean oils, 158 and monounsaturated fat can range up to 18% in recognition of the fact that replacement of saturated fat with monounsaturates leads to a smaller decrease in HDL-C than replacement with carbohydrate (especially important with low HDL-C levels). Two servings per week of fish have been shown to reduce the risk of sudden death and death due to CHD in adults. 159 Additional recommendations that can lower risk for CHD and improve the lipid profile include consumption of lean meats and fat-free milk, elimination of hydrogenated or partially hydrogenated fats, elimination of trans -fat, and reduction of cholesterol to less than 300 mg/day. 159 Alcohol restriction is important in those with TG levels > 500 mg/dL.

Recommended diets should be calorie restricted to promote



weight loss because even moderate weight reduction can lower levels of TGs. $^{\rm 111}$ In a meta-analysis, $^{\rm 160}$ TG levels

TABLE 15—5	Pharmacolog	ic Treatment of	Hypertriglyceric	lemia			0.1	
Drug Class	Decrease in TG (%)	Decrease in LDL-C (%)	Increase in HDL-C (%)	Maintenance Regimen	Contraindications	Side Effects	Selective Decrease in Small, Dense LDL-C	Selective Increase in HDL ₂ Cholesterol
Nicotinic acid, extended release	17-26	10-25	15-35	1500 Hypersensitivity, hepatic 2000 mg once dysfunction a day at bedtime	Flushing, pruritus, nausea hepatitis (at higher doses), activation of migraine (rare)	Yes	Yes	
Fibrate: gemfibrozil	35-50	No change to slight increase	15-25	Gemfibrozil, 600 mg twice a day	Hypersensitivity, hepatic dysfunction, end- stage renal disease	Myositis, cholelithiasis	Yes	No
Fibrate: fenofibrate	41-53	6-20	18-33	Fenofibrate, 145 mg once a day	Hypersensitivity, hepatic dysfunction, end- stage renal disease	Myositis, cholelithiasis	Yes	No
Statins	5	20-60	5-10	Multiple agents	Hypersensitivity, pregnancy, breast feeding	Myalgia, influenzalike syndrome, rhabdomyolysis (rare), weakness	No	No
Nicotinic acid and statin	20-35	30-65	20-45	Same as for individual agents	Same as for individual agents	Same as for individual agents	Yes	Yes
Statin and fibrate	35-55	20-60	15-30	Same as for individual agents	Same as for individual agents	Same as for individual agents	Yes	No
Cholesterol absorption inhibitors	No change	17	No change	Ezetimibe, 10 mg daily	Hepatic dysfunction	Hepatitis	No	No

decreased 0.015 mmol/L for each kilogram of weight lost. Small amounts of weight loss (5.3%) have been shown to result in approximately a 22% decrease in TG levels, a 9% increase in HDL-C, and a 40% decrease in small, dense LDL particles. 161,162 HL, which removes TG, can metabolize HDL to a smaller size, which has a higher FCR, thus lowering levels of HDL. In older men, reduction of intra-abdominal fat stores has resulted in lower levels of HL activity and thus higher levels of HDL-C. 161 In the Women On the Move through Activity and Nutrition (WOMAN) trial, lifestyle changes reduced the number of LDL particles as well. 163 In addition to improved lipid profiles, lifestyle interventions can lower the frequency of angina 164 and the need for interventions in those with CHD. 165 In the Clinical Outcomes Utilizing Revascularization and Aggressive Drug Evaluation (COURAGE) trial, two thirds of the patients with stable angina and objective evidence of myocardial ischemia in the medical treatment arm (which included a team approach with physicians, dietitians, nurses, and exercise specialists to make lifestyle changes) did well without invasive coronary intervention. 165

Current American Heart Association guidelines for exercise recommend moderate-intensity aerobic exercise for a minimum of 30 minutes for 5 to 7 days or vigorous-intensity aerobic exercise for a minimum of 20 minutes for 3 days a week. 166 Even in the absence of weight loss, aerobic exercise has been shown to modestly reduce TG levels in a dose dependent fashion. 167 Exercise can reduce levels of small, dense LDL particles, increase LDL particle size, raise HDL-C levels, and lower levels of TG. 168 A meta-analysis of 25 studies of walking has also shown an improvement in the lipid profile. ¹⁶⁹ In a comparison of four types of exercise varying in intensity and frequency, the hardintensity-high-frequency exercise was the only one to

significantly improve HDL-C (mean increase, 3.9%; P < 0.03). ¹⁷⁰ Initial weight loss and maintenance of weight loss can be

difficult in clinical practice; however, Welty and coworkers 162 described a successful office-based lifestyle program for modification of lipids and blood pressure that also resulted in weight loss. The program includes a dietitian at the initial visit and at least one follow-up visit (fully reimbursable to dietitian with separate co-pay). The diet described before was recommended with additional recommendations of low sodium, fiber > 25 g/day for women and 38 g/day for men, low-fat dairy products, fruits and vegetables, oils high in alpha-linolenic acid (canola and flaxseed oils), and plant sterols or stanols to 2 g/day, all geared towards lowering of TG, LDL-C, and blood pressure and raising of HDL-C. Behav ioral counseling to decrease portion size included (1) eating a high-fiber breakfast daily, (2) drinking a full glass of water before each meal and with high-fiber foods to induce satiety, (3) putting the fork down between each bite and taking a sip of water or stopping a meal halfway through for 15 to 20 minutes (time for satiety signal to reach brain) geared to slowing down eating, (4) decreasing portion size by 100 calories per day (equates to 10-pound weight loss in 1 year), and (5) taking half of a restaurant portion home. Exercise prescriptions were given and included increasing 5 minutes per session weekly to a goal of 30 minutes daily. Maximum weight lost was an average of 5.6% (10.8 pounds) at a mean follow-up of 1.75 years. Sixty-four (81%) of these patients maintained significant weight loss (average weight loss, 5.3%) at a mean follow-up of 2.6 years. Those maintaining weight loss exercised significantly more. 162 Average decreases in LDL-C and TG were 9.3% and 34%, respectively, and

244 average increase in HDL-C was 9.6%. ¹⁶¹ Diastolic blood pressure - fibrates reduces the risk of CHD events in patients with high TGs and maintain weight loss.

TREATMENT: **MEDICATIONS** TO **TREAT HYPERTRIGLYCERIDEMIA**

There are currently four major medications to treat elevated TG concentration: statins, fibrates, niacin, and fish oil. Table 15-5 summarizes average changes in lipid levels for these drugs. The following sections summarize the role of each drug in lowering TG concentration and the clinical trial evidence supporting its use.

Statins

Once the TG level is < 500 mg/dL, or if the TG level is 200 to 499 15 mg/dL, if the LDL-C level remains above goal, statins are the drug of first choice for LDL lowering because of their reduction in with baseline TG levels greater than 2.8 mmol/L had dosedependent reductions in TGs of 22% to 45%, with atorvastatin and rosuvastatin being the most effective. As noted earlier, in a 140 mg/dL, and TG < 300 mg/dL were randomized to gemfi brozil consensus statement, statins remained the first-line therapy for or placebo. Gemfibrozil raised HDL-C by 6% (34 mg/dL versus 32 moderate hypertriglyceridemia, and other TG-lowering drugs, such mg/dL), lowered TGs by 31% (115 mg/dL versus 166 mg/dL), and as fibrates and niacin, were suggested for high-risk patients. 145

the progression of coronary atherosclerosis in those with an LPL reduction in the primary endpoint of nonfatal MI or death due to mutation. Heterozygosity for the Asp9Asn mutation in the LPL gene CHD (17.3% versus 21.7% for placebo; relative risk reduction, 22%; (aspartic acid to an asparagine residue at position 9) causes mild P = 0.006). ¹⁸⁶ All-cause mortality was lower by a nonsignificant 11%. defects in LPL activity leading to elevations in serum TGs and a When stratified by diabetic status, diabetics had a significant 32% reduction in HDL-C. 1,173 In the Regression Growth Evaluation Statin reduction in the combined primary endpoint (P = 0.004), death due Study (REGRESS), subjects with this mutation were more likely to to CHD (P = 0.02), and stroke (P = 0.046) compared with those have a family history of cardiovascular disease (P = 0.03), a lower without diabetes (P = 0.07, P = 0.88, and P = 0.67, respectively). HDL-C level (P = 0.01), a trend toward higher TG and LDL-C levels, Eight-year follow-up in VA-HIT showed a 41% reduction in CHD and progression of coronary atherosclerosis in the placebo group death (P = 0.02) and a 26% reduction in death in those with diabetes compared with noncarriers. Therapy with pravastatin attenuated or hyperinsulinemia at baseline. There fore, diabetics were more the rate of progression of atherosclerosis in those carrying the likely than those without diabetes to benefit (see Table 15-6), mutation. 174

Fibrates

proliferator-activated receptor a (PPAR a), fibrates decrease the apo C-III concentration, which increases LPL activity, thus increasing glucuronidation; therefore, fenofibrate is safer to use in combination VLDL clearance and reducing the plasma TG concentration by 30% to 50% (see Table 15-5). 175 Fibrates also increase fatty acid oxidation cardiovascular disease outcomes. In the Diabetes Atherosclerosis in liver and muscle and reduce the rate of lipogenesis in the liver, thereby reducing hepatic secretion of VLDL TGs. 175 PPAR a mediated transcription by fibrates also stabilizes apo AI mRNA Those taking fenofibrate had less progression of coronary artery transcripts, 176-178 leading to secretion of more apo AI-containing disease compared with those randomized to placebo. 187 Part of the HDL particles , thus increasing HDL production and raising the benefit appeared to be related to an increase in the size of LDL HDL-C level by 15% to 30%, depending on baseline HDL-C level (see Table 15-5). HDL-C levels are also raised by the transfer of endpoints, the rate of all-cause mortality was 2.9% in the treatment surface apo AI released during the enhanced catabolism of VLDL to HDL particles. The effect of fibrates on LDL-C ranges from around a 10% reduction to no change or a slight increase and an increase in (FIELD) study was a 5-year study of 9795 patients with LDL particle size. The accelerated clearance of TG-rich lipoproteins reduces CETP-mediated exchange, which results in less TG incorporation into LDL particles and thus limits formation of small, dense LDL and favors the formation of large, cholesterol-enriched LDL particles 125,179 that bind more efficiently to the LDL receptor than small LDL particles do. Thus, there is an overall reduction in LDL particle concentration despite a minimal change in LDL-C level. 179 Fibrates may also have anti-inflammatory and antiatherogenic properties due to activation of PPAR a . 180

Several studies have shown that monotherapy with various

decreased from a mean of 79 to 75 mm Hg (P = 0.003). Therefore, and low HDL-C, especially in patients with diabetes mellitus or having a dietitian counsel patients concurrently with a physician characteristics of the metabolic syndrome (summarized in Table 15and using the simple techniques described here may help achieve 6). 181 In the Bezafibrate Infarction Prevention (BIP) trial, 3090 subjects with prior MI and HDL-C < 45 mg/dL, TG < 300 mg/dL, and LDL-C < 180 mg/dL were randomized to bezafibrate 400 mg daily or placebo. Increasing serum TG concentrations were associated with an increase in mortality. 182 Although bezafibrate decreased levels of LDL-C by 6% and TGs by 21% and raised levels of HDL-C by 18%, there was no significant difference in the primary endpoint (fatal and nonfatal MI and sudden death) in either group. However, those patients with TG levels > 200 mg/dL at baseline who had an increase in HDL of at least 5 mg/dL or a reduction in TG by > 28 mg/dL had a significant 40% (13% versus 22.3% at 6.2 years; P = 0.02) reduction in the primary endpoint. ¹⁸³

Gemfibrozil has been studied in several trials. In the Hel sinki Heart Study (HHS), a 5-year primary prevention trial, dyslipidemic men without CHD randomized to gemfibrozil had a significant 34% reduction in the incidence of fatal and nonfatal MI and cardiac death. 184 All-cause mortality did not differ significantly between the cardiovascular events, stroke, coronary mortality , and total treatment group (21.9%) and the placebo group (20.7%). Those with mortality without an increase in noncardiovascular mortality. 171 In TG levels > 200 mg/ dL and LDL-C/HDL-C ratio > 5.0 had the an analysis of the efficacy of statins in lowering TG levels, 172 patients greatest risk of CHD, and those in this group treated with gemfibrozil had a 71% event reduction (see Table 15-6). 185

In VA-HIT, men with CHD and HDL-C < 40 mg/dL, LDL-C < had no significant effect on LDL-C (mean 113 mg/dL in both An angiographic trial has shown that statin treatment attenuates groups). Those treated with gemfibrozil had a significant 22% possibly because of improvement in insulin resistance through PPAR a activation.

Gemfibrozil is more likely to cause myositis in combination with a statin by raising plasma levels of statins 1.9- to 5.7-fold through By activating the nuclear transcription factor peroxisome inhibition of hepatic glucuronidation, which is necessary for renal excretion of lipophilic statins. 111 Fenofibrate does not inhibit with a statin. Several trials have studied the effect of fenofibrate on Intervention Study (DAIS), 731 subjects with diabetes were randomized to micronized fenofibrate (200 mg/dL) or placebo. particles. Although the study was not powered for clinical group and 4.3% in the placebo group.

The Fenofibrate Intervention and Event Lowering in Diabetes

	the Metabol	ic Syndrome				
Study	Treatment	Patients	Endpoint	Absolute Risk Reduction (%)	Relative	P Value
Primary P	revention					
HHS	Gemfibrozil	4081 men with no prior CVD	Nonfatal MI + CAD death	13.8	34	< 0.02
	1200 mg/day	1146 (28% with MS) 292*	Nonfatal MI + CAD death Nonfatal MI + CAD death	27.2 9.1	71 71	<0.0005 0.004
FIELD	Fenofibrate 200 mg/day	9795 men and women with type 2 diabetes (some with CAD)	Nonfatal MI + CAD death	0.7	11	NS
		7664 without CHD	Total CVD events Nonfatal MI + CAD death	3.2 1.9	11 19	0.035 0.004
		8183 with MS 314 with MS and marked dyslipidemia-	Total CVD events Total CVD events	1.4 4.3	11 27	0.052 0.005
Secondary	y Prevention					
VA-HIT	Gemfibrozil 1200 mg/day	2531 men with CAD	Nonfatal MI, CAD death + stroke	5.6	24	<0.001
		769 (30%) with diabetes	Nonfatal MI, CAD death + stroke	10.0	32	0.004
BIP	Bezafibrate 400 mg/day	3090 men and women with previous MI	Nonfatal MI + CAD death	1.4	9.4	NS
	3.11	1470 (48%) with MS-		4.3	25	0.03

Monotherapy in Patients with and without Diabetes or

type 2 diabetes (78% without CVD) with TC from 116 to 251 mg/dL fibrates. Fibrates did not significantly reduce the odds of plus either a TC/HDL-C ratio of at least 4.0 or a TG concentration of cardiovascular mortality (P = 0.68), fatal MI (P = 0.76), or stroke (P = 0.68), fatal MI (P = 0.76), or stroke (P = 0.68), fatal MI (P = 0.76), or stroke (P = 0.68), fatal MI (P = 0.76), or stroke (P = 0.68), fatal MI (P = 0.76), or stroke (P = 0.68), fatal MI (P = 0.76), or stroke (P = 0.68), fatal MI (P = 0.76), or stroke (P = 0.76), or stroke (P = 0.76), or stroke (P = 0.76), fatal MI (P = 0.76), or stroke (P = 0.76), or stroke (P = 0.76), or stroke (P = 0.76), fatal MI (P = 0.76), or stroke (P = 0.76), or stroke (P = 0.76), fatal MI (P = 0.76), or stroke (P = 0.76), or stroke (P = 0.76), fatal MI (P = 0.76), or stroke (P = 0.76), or stroke (P = 0.76), fatal MI (P = 0.76), fatal MI (P = 0.76), or stroke (P = 0.76), fatal MI 89 to 443 mg/dL. Those randomized to fenofibrate 200 mg/day had = 0.56), and these results did not change with the exclusion of a nonsignificant 11% reduction (P = 0.16) in the primary endpoint of clofibrate trials. However, fibrates significantly reduced the odds of first MI or CHD death compared with those receiving placebo (see Table 15-6). ¹⁸⁸ The initiation of statin therapy in many patients reduction (pooled odds ratio, 0.75; P < 0.00001) with exclusion of stating the trial unfortunately approximate the trial union of a significant reduction of 25% (P = 0.014) in the primary endpoint. randomized controlled trials reported a 13% increase in risk for For the overall group, significant reductions were observed in the secondary outcomes of nonfatal MI (HR, 0.76; 95% CI, 0.62-0.94), nine of these trials used clofibrate, and FIELD was not included. 171 total cardiovascular disease events (HR, 0.89; 95% CI, 0.80-0.99), When outcome in fibrate trials is examined for patients with coronary revascularization (HR, 0.79; 95% CI, 0.68-0.93), and all diabetes or metabolic syndrome, a greater benefit is seen in both revascularizations (HR, 0.80; 95% CI, 0.70-0.92). Total mortality, primary prevention (HHS and FIELD) and secondary prevention receiving placebo. In a post hoc subgroup analysis in FIELD, those death (P = 0.0001). ¹⁹¹ Therefore, fibrates may have greater benefit in this group, in whom a 27% relative risk reduction in total CVD events occurred (HR, 0.73; 95% CI, 0.58-0.91; P = 0.005; number needed to treat = 23) compared with 6% in all others (HR, 0.94; 95% vascular Risk in Diabetes (ACCORD) study was designed to CI, 0.83-1.06; P = 0.321; number needed to treat = 143). 189

TABLE 15- -6 Cardiovascular Risk Reduction with Fibrate

randomized placebo-controlled trials including the FIELD trial, fibrates significantly reduced plasma TC and TG levels by about 8% and 30%, respectively, and raised HDL-C levels by about 9% compared with placebo. 190 The odds of all-cause mortality were higher (P = 0.08) and the odds of noncardio-vascular mortality were significantly higher (P = 0.004) with the use of fibrates. However, after exclusion of trials that used clofibrate as the study drug (because of withdrawal of clofibrate from the market more than 25 years ago due to an increase in cholangiocarcinoma and other gastrointestinal cancers), neither all-cause mortality (pooled odds ratio, 1.04; P = 0.44) nor noncardiovascular mortality (pooled odds ratio, 1.08; P = 0.20) was significantly increased with the use of

during the trial, unfortunately, confounded the results. The statin clofibrate trials. The odds for development of cancer were not 🕏 drop-in rate was higher in those with prior CVD compared with significantly higher with the use of fibrates (P = 0.98), nor were the those without CVD. Analysis of patients with no prior CVD showed odds of cancer-related death (P = 0.17). Another meta-analysis of 17 those without CVD. Analysis of patients with no prior CVD showed odds of cancer-related death (P = 0.17). Another meta-analysis of 17

however, was increased by a nonsignificant 11% in those receiving (BIP and VA-HIT) in those with diabetes or metabolic syndrome fenofibrate (HR, 1.11; 95% CI, 0.95-1.29), and pancreatitis and compared with those without (see Table 15-6). Patients in the pulmonary embolism were significantly higher in those receiving highest tertile of body mass index and TG level in follow-up in HHS fenofibrate (P = 0.03 and 0.022, respectively) compared with those had a reduction of 33% in risk of death (P = 0.03) and 71% in CHD with metabolic syndrome and marked dyslipidemia (TG > 2.3 cardioprotective effects in patients with diabetes mellitus or features mmol/I [204 mg/dL] and low HDL-C) had the highest risk (17.8%) of the metabolic syndrome because of improvement in insulin for development of CVD during 5 years. Fenofibrate had the greatest resistance through PPAR agonist activation. These prior studies did determine whether fibrate therapy provided additional benefit to In a meta-analysis of 36,489 patients from 10 published statin therapy in subjects with type 2 diabetes. 192 In the

^{*}Patients with TG >204 mg/dL and an LDL/HDL >5 (may or may not have had diabetes mellitus or the metabolic syndrome).

Defined as elevated triglycerides (>204 mg/dL [>2.3 mmol/L]) and HDL <1.03 mmol/L for men and <1.29 mmol/L for women.

Defined as elevated triglycerides (>204 mg/dL [1 mg/dL = 0.0113 mmol/L]), low HDL-C (<42 mg/dL [1 mg/dL = 0.02586 mmol/L]), body mass index >26, and blood glucose >5.5

BIP, Bezafibrate Infarction Prevention; CAD, coronary artery disease; CVD, cardiovascular disease; FIELD, Fenofibrate Intervention and Event Lowering in Diabetes; HHS, Helsinki Heart Study; MI, myocardial infarction; MS, metabolic syndrome; NS, not significant; VA-HIT, Veterans Affairs High-Density Lipoprotein Intervention Trial.

246 trial, 10,251 patients with type 2 diabetes were randomized to either intensive or standard glycemic control, with 5518 also randomly assigned in a 2 x 2 factorial design to either simvastatin plus fenofibrate or simvastatin plus placebo. Fenofibrate plus simvastatin did not reduce the primary outcome (fatal CVD events, nonfatal MI, or nonfatal stroke) compared with simvastatin alone in the total group (2.2% versus 2.4%, respectively; HR, 0.92; 95% CI, 0.79-1.08; *P* = 0.32). However, those with TG levels > 204 mg/dL (> 2.3 mmol/L) (TG level was lowered an average of 35%) and HDL level < 34 mg/dL (HDL-C increased an average of 12.9%) randomized to fenofibrate had a 28% reduction in relative risk (12.4% event rate compared with 17.3% with placebo [P = 0.057]) compared with rates of 10.1% in both study groups for all other subjects. Therefore, although the routine use of combination therapy with fenofibrate and simvastatin did not improve outcome in subjects with type 2 diabetes, the ACCORD subgroup results suggest that the addition of fenofibrate to a statin provides additional benefit in type 2 diabetics with high TG and low HDL-C levels. In contrast to the adverse outcomes in FIELD, no increase in total mortality, pulmonary embolus, or

of cyclosporine, leading to an approximately 30% reduction in longer recommended for this purpose. ²⁰⁵ plasma level ¹⁸⁸; therefore, gemfibrozil is the preferred choice in transplant recipients. Fenofibrate lowers levels of fibrinogen and uric acid, whereas gemfibrozil has no effect.

Niacin

acid can increase blood glucose concentration, but doses of 750 (Lovaza) do not. to 2000 mg/day affect glucose only modestly. In the Arterial to 125 mg/dL, respectively.

events in the HDL-Atherosclerosis Treatment Study (HATS), Familial Atherosclerosis Treatment Study (FATS), and Cholesterol-Lowering Atherosclerosis Study (CLAS). 197-200 In the HATS trial, 160 patients with clinical and angiographic evidence of CHD, HDL-C < 35 mg/dL, and LDL-C < 145 mg/dL were randomized to one of four regimens: simvastatin plus niacin; antioxidants; simvastatin plus niacin plus

antioxidants; or placebo. Simvastatin plus niacin was associated with a 42% reduction in LDL-C and 26% increase in HDL-C and regression of proximal coronary plaque on coronary angiography (0.4% on average versus baseline) compared with a mean progression of 3.9% on placebo. 198 Those in the sim vastatin plus niacin group were also significantly less likely to experience a CVD event, death, MI, stroke, or revascularization, although the numbers were small. Although niacin use is limited by flushing, taking an aspirin (325 mg) 30 minutes before extended-release niacin at bedtime, taking it with a snack, and avoiding alcohol and spicy foods limit flushing. Education of patients about the fact that flushing is also an indication that the drug is working and that they are properly responding to it is key to maximizing adherence. Flushing may also be transient and may decrease with time in many cases. The Atherothrombosis Intervention in Metabolic Syndrome with Low HDL-C/High Triglyceride and Impact on Global Health Outcomes (AIM-HIGH) study is examining the use of niacin plus statins and should clarify the role of niacin in the treatment of hypertriglyceridemia. 201

Extended-release niacin can lower lipoprotein(a), an important pancreatitis was observed in the fenofibrate group compared atherogenic lipoprotein, approximately 20% at doses of 2 g daily. 202 with placebo in ACCORD; however, there was a trend Immediate-release niacin can lower lipoprotein(a) 38% at 4 g daily. towards harm in women and benefit in men in the total group. 203 The only other medication that can lower lipoprotein(a) and is Both gemfibrozil and fenofibrate interfere with metabolism available in the United States is estrogen replacement therapy, of warfarin; therefore, warfarin dose should be decreased which lowered lipoprotein(a) 17% to 23% in the Postmenopausal approximately 30%, with close monitoring of the international Estrogen/Progestin Intervention trial 204; however, because of its normalized ratio (INR). 193 Fenofibrate increases the clearance potentially harmful effects, estrogen replacement therapy is no

Omega-3 Fatty Acids

Fish oil supplements (>3 g/day of eicosapentaenoic acid [EPA] and docosahexaenoic acid [DHA], the active omega-3 fatty acids) can lower TG level by 50% or more by reducing VLDL production. ²⁰⁶⁻²⁰⁸ The extended-release form of niacin reduces TGs by 17% to However, use can be limited by reflux and regurgitation of a "fishy 26% and raises HDL-C 15% to 35% with a dose of 2000 taste." Placement of tablets in the refrigerator or freezer can mg/day (see Table 15-5). 148 A 15% reduction in risk of MI minimize this side effect. An enteric-coated capsule is also on the occurred in 8341 men with prior MI and hypercholesterolemia market that decreases the fish odor problem. Table 15-7 outlines the who were randomized to regular-release nicotinic acid 3 g number of tablets and approximate cost for each over-the-counter daily or placebo and observed for 6.5 years in the Coronary fish oil preparation to obtain 1 g EPA plus DHA daily. We aim for at Drug Project. 194 Moreover, total mortality was significantly least 2.4 g EPA plus DHA daily in our patients with reduced 11% (P = 0.0004) and CHD mortality was reduced hypertriglyceridemia. Over-the-counter fish oil capsules have some 12% (P < 0.05) at 15-year follow-up. 194 High doses of nicotinic cholesteryl ester, whereas prescription ethyl esters of omega-3

Omega-3 fatty acids can be an alternative to fibrates or niacin, Disease Multiple Intervention Trial (ADMIT), 2 to 3 g/day of although they can also be used in combination with statins, fibrates, immediate-release niacin decreased TG by 23% and raised or niacin. 111 In addition to effectively lowering TGs, omega-3 fatty HDL by 29% without significant worsening of glycemic acids may provide clinical benefit in patients with recent MI and control. 195 In the Assessment of Diabetes Control and may have antithrombotic and anti-inflammatory effects. 209 There Evaluation of the Efficacy of Niaspan Trial (ADVENT) in 97 are three large trials of fish oil. The largest trial with omega-3 fatty patients with type 2 diabetes, 1 to 1.5 g/day of extended- acids was the Gruppo Italiano per lo Studio della Sopravvivenza release niacin resulted in no significant difference in mean nell-Infarto Miocardico (GISSI) Prevenzione trial, which showed a fasting glucose levels during a 16-week period, although 20% reduction in the combined endpoint of death, nonfatal MI, and additional hypoglycemic treatment was required in some stroke with 1 g/day of EPA plus DHA. 210 Omega-3 fatty acids also patients receiving niacin. 196 Six-year follow-up in the had significant antiarrhythmic effects. 210,211 In the Japan Coronary Drug Project showed that those with a baseline Eicosapentaenoic Acid Lipid Intervention Study (JELIS), the glucose concentration of > 126 mg/dL had the greatest combination of open-label low-dose statin (pravastatin, 10 to 20 mg, reduction in risk of recurrent MI (57%) compared with 30%, or simvastatin, 5 to 10 mg, daily) with open-label eicosapentaenoic 24%, and 25% reductions in those with baseline fasting plasma acid (1800 mg daily) reduced incident and recurrent CHD events by glucose concentration < 95 mg/dL, 95 to 104 mg/dL, and 105 19% (P = 0.011) compared with statin monotherapy. ²¹² There was no effect on cardiac or total mortality. LDL-C levels were reduced Niacin in combination with simvastatin lowered CVD by 25% in both groups, whereas TG levels were

Brand or Type	No. of Capsules for — 1 g EPA + DHA	Cost for 30 Days*	Notes
GNC Triple Strength Fish Oil (647 mg EPA, 253 mg DHA per softgel)	'	\$10	60 caps/\$19.99
Vitamin World Triple Strength Omega-3 Fish Oil 1360 mg (EPA + DHA 950 mg per softgel) Same as Puritan's Pride	1	\$11	Size: slightly > 1 inch 60 caps/\$21.99 Size: 1 inch
Lovaza 1000 mg (prescription only) (approx. 465 mg EPA and 375 mg DHA)	1	\$40-\$60	Insurance may not cover or may have the highest co-pay
Frader Darwin's Omega 3 Fatty Acids 1100 mg (300 mg EPA, 200 mg DHA per softgel)	2	\$5	Trader Joe's brand 90 caps/ \$7.99 Size: 1 inch
Frader Darwin's Omega 3 Fatty Acids - Odorless 1200 mg 400 mg EPA, 200 mg DHA per softgel)	2	\$6	Trader Joe's brand 90 caps/\$8.99 Size: 1 inch
	2	\$6	
(irkland Signature Fish Oil Concentrate 1200 mg enteric coated 410 mg EPA, 270 mg DHA per softgel)			Costco's brand 180 head/\$16.99 Size: slightly > 1 inch
Walgreens Finest enteric-coated One-Per-Day Omega 3 Fish Oil 1200 mg 410 mg EPA, 274 mg DHA per softgel)	2	\$10	60 caps/\$9.99 Size: 1 inch (enteric coated)
Nature Made Fish Oil Double Strength 1200 mg (336 mg EPA, 276 mg DHA per softgel)	2	\$17	60 caps/\$16.99 Size: ?
/itamin World Premium Mini Gels Omega-3 Fish Oil (EPA + DHA 450 mg per softgel)	2	\$18	60 caps/\$17.99 Size: 34 inches
the country per conger,	2	\$28	
Nordic Naturals Ultimate Omega, Highly Concentrated 325 mg EPA, 225 mg DHA per softgel)			IFOS · Size: 1 inch (lemon flavor)
Puritan's Pride Natural Omega-3 1000 mg (300 mg EPA + DHA per softgel) Same as Vitamin World	3	\$7	100 caps/\$10.99 Size: 1 inch
Whole Foods molecularly distilled Omega-3 1000 mg (180 mg EPA, 120 mg DHA per softgel)	3	\$10	90 caps/\$9.99
			Size: 1 inch (lemon flavor)
Gels and liquids Coromega Omega-3 Supplement (350 mg EPA, 230 mg DHA per squeeze packet)	2 packs	\$20-\$40	
Nordic Naturals Omega-3 Liquid, lemon taste (165 mg EPA, 110 mg DHA per mL)	— 3/4 tsp	\$12	Flavors: orange, orange-chocolate, lemon-lim 8oz/\$24.95 Keep refrigerated, spoils easily

^{*}Cost data were obtained from locations in the Boston, Mass, area in October 2009.

Most commercial brands have more than one formulation. The less expensive options are typically not as concentrated and tend to vary the EPA and DHA content with other fish oil components. Testing is not performed on all over-the-counter supplements sold; however, the majority of fish oil supplements have been shown to be free of contaminants, such as mercury. Capsules that have a strong odor may be spoiled and should not be used. To keep supplements fresh, store them in the refrigerator or freezer. Cod liver oil, which may contain excess vitamin A, is not recommended.

unchanged. In the third trial, patients with chronic heart failure had significant reductions in coronary events and cardiac and total mortality. 213

In subjects with persistent hypertriglyceridemia, omega-3 fatty acid prescription at 4 g/day plus simvastatin 40 mg provided significant additional improvement in reducing levels of non-HDL-C (- 9% versus - 2.2%, respectively; P < 0.001), TG (29.5% versus 6.3%, respectively; P < 0.001), and other lipid and lipoprotein parameters to a greater extent than simvastatin alone. ²¹⁴ Whether other combinations involving TG-lowering therapies are also clinically superior to statin monotherapy is a subject of intense investigation. Ongoing trials include the addition of niacin to statin, which

is being investigated in AIM-HIGH 201 and HPS2-THRIVE. 215 The results of these trials are expected to help shape evidence-based guidelines to optimize management of dyslipidemia , including elevated levels of TG.

Acid Sequestrants Balls

Because of increases in VLDL synthesis and increases in TG levels, bile acid sequestrants should not be used unless TG levels are normal. Furthermore, they are normally reserved for treatment of elevated LDL-C when statins cannot be tolerated or are contraindicated or in combination with statins to achieve greater LDL-C lowering.

^{*}ConsumerLab tested; IFOS, International Fish Oil Standards.

of therapy when TG levels are in the borderline-high range (150 to 199 mg/dL) because apo B-containing lipoproteins can also be elevated in this setting. Lifestyle changes — exercise and diet with an emphasis on weight loss — are first-line therapy in lowering TG levels, especially in patients with metabolic syndrome or type 2 diabetes. Statins, fibrates, niacin, and omega-3 fatty acids can be used to lower TG levels to reach the non-HDL-C goal if lifestyle changes are insufficient to reach the goal. On the basis of the ACCORD trial results, fenofibrate can also be considered in addition to a statin in type 2 diabetics with marked dyslipidemia (TG > 204 mg/dL and HDL-C < 34 mg/dL) to lower the risk of CVD events. $^{\rm 192}$

Hypertriglycgidemia is associated with other lipid -

abnormalities that predispose to atherosclerosis, including low

levels of HDL-C, the presence of small, dense LDL particles and

atherogenic TG-rich lipoprotein remnants, and insulin

resistance. The evidence cited in this chapter suggests that non-

HDL-C (TC - HDL-C) may predict risk of CHD and future

cardiovascular events better than LDL-C because of the fact that

non-HDL-C includes all apo B-containing lipoproteins, all of

which are atherogenic. Therefore, NCEP recommends

calculation of non-HDL-C when the TG level is > 200 mg/dL,

with the goal being 30 mg/dL higher than the LDL-C goal. It is

also not unreasonable to consider non-HDL-C a secondary target

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Diet and Lifestyle Factors

CHAPTER 16

Nutritional Approaches for Cardiovascular Disease Prevention

Alison M. Hill, Kristina A. Harris, Alison M. Coates, and primary focus of CVD prevention is analytic Kris-Etherton

- pressure lowering and improvement of other cardiometabolic risk factors.
- Current dietary guidelines for chronic disease prevention emphasize an overall healthy dietary pattern based on foods rather than on targeting specific nutrients.
- Strategies used in clinical practice for CVD prevention, specifically LDL-C and blood pressure lowering, include the Therapeutic Lifestyle Changes (TLC) and Dietary Approaches to Stop Hypertension (DASH) diets.
- Dietary patterns for weight loss are most dependent on calorie control, but modification of the macronutrient profile may target cardiovascular risk factors, including blood pressure, lipids, and lipoproteins.
- Key components of hearthealthy dietary approaches include reduced intake of saturated fatty acids, trans-fatty acids, and cholesterol and increased intake of fruits and vegetables, whole grains. reduced- fat dairy, and hearthealthy protein (from plant and low-saturated fatty acid animal sources).

· Additional dietary components (soluble fiber, sterols and stanols, soy protein, and unsaturated fats) and supplements (fish oil and niacin) improve CVD risk status.

NUTRITIONAL GOALS FOR PREVENTION OF **CARDIOVASCULAR DISEASES**

The goal of cardiovascular disease (CVD) prevention is to decrease CVD morbidity and mortality through pharmacological and lifestyle (including dietary, behavioral, and physical) intervention. Reduction in major risk factors, which include total cholesterol and low-density lipoprotein cholesterol (LDL-C), systolic blood pressure, smoking prevalence, and physical inactivity, accounted for almost half of the decrease in coronary heart disease (CHD) mortality in the United States between 1980 and 2000. ¹ Healthy lifestyle practices that improve such modifiable risk factors are therefore imperative for CVD prevention, and unlike pharmacological treatments, lifestyle interventions are accessible to everyone and have similar efficacy in "slowing" chronic disease progression. ² Modest dietary changes across an extended period can produce tangible results, both physically and psychologically, and may have a significant financial benefit. For example, if every adult in the United States reduced total daily calorie intake by 100 kcal, approximately 71.2 million cases of

· Food-based recommendations and their associated tools, such as MyPyramid (developed by the US Department of Agriculture), assist consumers in implementing dietary guidelines and may assist clinicians in counseling patients.

overweight and obesity could be resolved, saving an estimated \$58 billion annually. 3

The most prominent risk factors for CVD are elevated serum total cholesterol and LDL-C, hypertension, diabetes, and cigarette smoking. 4,5 Randomized controlled trials have shown that lowering LDL-C and blood pressure, in particular, reduces the risk for CVD. 6,7 Statin drug trials demonstrate that for every 25 mg/dL lowering of serum LDL-C, there is a decrease in major vascular (-14 %) and coronary (-16 %) events. 8 A linear relationship also exists between hypertension and CVD risk, which doubles with every increment of 20/10 mm Hg in systolic blood pressure (SBP)/diastolic blood pressure (DBP). 6 Currently, the primary aim of nonpharmacological dietary interventions is a reduction in LDL-C, with a secondary aim of lipoprotein lowering non-high-density cholesterol and blood pressure. Other CVD risk factors, such as increased serum triglycerides (TG) and glucose, decreased high-density lipo protein cholesterol (HDL-C), increased hypertension. and circumference, which cluster as part of the metabolic syndrome, are also targets for intensive lifestyle therapy in an effort to optimize CVD prevention. 9 Evidence for nontraditional risk factors,

such as markers of thrombosis and inflammation, LDL-C particle size, and apolipoproteins, is emerging, 10 and additional research will establish their contribution to overall CVD risk.

Current dietary guidelines advocate a food-based approach for optimal health and chronic disease risk reduction. 11 Indeed, the American Heart Association (AHA) recommends a diet consistent with current guidelines to meet their 2020 Impact Goal to "improve the cardiovascular health of all Americans by 20% while reducing deaths from cardiovascular diseases and stroke by 20%." 12 The Dietary Guidelines for Americans, 2005 emphasize a balanced diet that incorporates a variety of foods that are consumed in moderation. There fore, although specific foods and nutrients have been identified as being more or less "healthful," it is the total dietary package that is most important. This chapter discusses the efficacy of a range of heart-healthy dietary patterns that have been scientifically evaluated for overall health promotion and CVD risk reduction. In particular, it reviews specific dietary patterns, such as the Therapeutic Lifestyle Changes (TLC) ⁹ and Dietary Approaches to Stop Hypertension (DASH) 13 diets, that are used in clinical practice for LDL-C and blood pressure lowering, respectively. Dietary strategies for weight loss are also reviewed. In evaluating dietary patterns, it is evident that a number of common foods exist between diets. The evidence supporting the benefits of 16 specific foods and supplements on cardiovascular risk factors, with a particular focus on reducing LDL-C and blood pressure, is Modified from Third Report of the National Cholesterol Education Program (NCEP) Expert discussed. The information presented in this chapter will assist clinicians in implementing dietary strategies to manage CVD risk Treatment Panel III). Final Report. Bethesda, MD, 2002, National Heart, Lung and Blood status in at-risk patients.

DIETARY	PATTERNS	TO	REDUCE
CARDIOVAS	CUI AR DISEASE	RISK	

Therapeutic Lifestyle Changes Diet

LDL-C has long been identified by the National Cholesterol Education Program (NCEP) as the primary target for CHD risk reduction. Recommendations by NCEP for testing and management of high blood cholesterol have been published in reports by the Expert Panel on Detection, Evaluation, and Treatment of High Blood Cholesterol in Adults (Adult Treatment Panel III). 9 The evidence base for such recommendations is primarily from randomized clinical trials that have demonstrated that lowering LDL-C substantially reduces the risk for CHD. Since the publication of the Adult Treatment Panel III (ATP III) guidelines, the results from several large-scale clinical trials have reinforced ATP III recommendations for therapeutic LDL-C lowering in high-risk and very-high-risk individuals. 14 ATP III recommends TLC for the clinical management of LDL-C in persons from all risk classifications (see Table 16-1 for summary of key components). 9 TLC may further reduce the incidence of CHD by modifying other cardiovascular risk factors beyond LDL-C. The Treatment Panel III). Final Report. Bethesda, MD, 2002, National Heart, Lung, and Blood primary focus of TLC is a reduction in saturated fatty acids (SFA) Institute. NIH publication 02-5215. to < 7% of total calories, trans -fatty acids to as low as possible, and cholesterol to < 200 mg/day. Guidelines also are provided for monounsaturated fatty acid (MUFA) and polyunsaturated fatty acid (PUFA) intake (up to 20% and 10% of total calories, 24% to 37% (Table 16-2). The magnitude of these effects has been respectively), carbohydrate intake (50% to 60% of total calories), established in well-controlled dietary intervention trials and protein intake (approximately 15% of total calories), and dietary further evaluated in free-living settings. The TLC dietary fiber intake (20 to 30 g/day). Additional therapeutic options for guidelines have been translated to food-based recommendations lowering LDL-C include viscous fiber (10 to 25 g/day) and plant that are consistent with the AHA Diet and Lifestyle stanols or sterols (2 g/day). Achieving or maintaining a healthy Recommendations 2006 (see Box 16-1, discussed later in this body weight (by calorie manipulation) and participating in regular chapter). Numerous dietary patterns that meet current nutri ent physical activity (enough moderate exercise to expend at least 200 recommendations for CVD risk reduction have been evaluated and kcal/day) are also essential TLC features. The combination of TLC shown to be efficacious relative to improving major risk factors for with other LDL-C-lowering options may collectively reduce LDL- CVD. Several of these dietary patterns are discussed on the C levels by

Component	Recommendation
LDL-raising nutrients Saturated fatty acids Cholesterol	< 7% of total calories < 200 mg/day
Therapeutic options for LDL lowering Increased viscous (soluble) fiber Plant stanols and sterols	10-25 g/day 2 g/day
Energy intake	
	Adjust calorie intake to achieve a healthy body weight or to prevent weight gain
Physical activity	
	Participate in enough moderate exercise to expend at least 200 kcal/day
Macronutrients Monounsaturated fatty acids Polyunsaturated fatty acids Total fat Carbohydrates Protein Dietary fiber	Up to 20% of total calories Up to 10% of total calories 25%-35% of total calories 50%-60% of total calories Approximately 15% of total calories 20-30 g/day

Panel on Detection, Evaluation, and Treatment of High Blood Cholesterol in Adults (Adult Institute. NIH publication 02-5215.

TABLE 16—2 Cumulative E	ffect of Cholesterol-Low	ering Strategies on LDL-
Dietary Component	Dietary Change	Approximate LDL Reduction
Saturated fat	< 7% of calories	8%-10%
Trans- fat	< 1% of calories	1%-2%
Dietary cholesterol	< 200 mg/day	3%-5%
Weight loss	Lose 10 pounds	5%-8%
Soy protein		3%-5%
Other LDL-lowering options Viscose fiber Plant sterol and stanol esters	5 to 10 g/day 2 g/day	3%-5% 6%-15%
Estimated cumulatives		24%-37%

Modified from Third Report of the National Cholesterol Education Program (NCEP) Expert Panel on Detection, Evaluation, and Treatment of High Blood Cholesterol in Adults (Adult

following pages.

Diet Portfolio

The portfolio diet incorporates four key cholesterol-lowering strategies, plant sterols (1.0 g/1000 calories), viscous fiber (8.2 g/1000 calories), soy protein (22.7 g/1000 calories), and almonds (14 g/1000 calories), within a reduced total fat diet (< 30% of total interventions, such as the portfolio diet, that address multiple CVD calories). 15 The portfolio diet is largely vegetarian, and SFA is risk factors, both traditional and emerging, limited to < 7% of total calories and cholesterol to < 200 mg/day. Sources of viscous fiber include eggplant and okra, oat bran, barley grains, and psyllium. Soy protein is obtained from soy milk and other soy products (such as soy burgers and sausages), and beans, chick peas, and lentils provide additional vegetable protein.

In a controlled 1-month dietary intervention with hyper lipidemic individuals, the portfolio diet reduced LDL-C by 29%, which is rich in fruits and vegetables (8 to 10 servings/day) and compared with an 8% reduction achieved by a low SFA (<7% of low-fat dairy products (2 or 3 servings/day), includes whole total calories) and cholesterol (< 200 mg/day) diet alone (also grains, legumes, fish, and poultry and is limited in added sugars known as a Step II diet). 15 Å subsequent trial compared the and fats. It is high in dietary fiber I (-30 g/day), magnesium, effectiveness of the portfolio diet on LDL-C lowering with statin potassium, and calcium and low in total fat (27% of total calories), therapy. ¹⁶ In this study, subjects completed three 1-month SFA (< 7% of total calories), and cholesterol (150 mg/day). In this intervention treatments in a randomized crossover design: a very study, 459 adults with 16 mildly elevated blood pressure (SBP < 160 low saturated fat diet (control or Step II diet), a control diet plus 20 mm Hg and DBP 80 to 95 mm Hg) were randomized to consume a mg lovastatin (statin), and the portfolio diet. LDL-C was reduced Western diet (control diet; 48% carbohydrate, 15% protein, 37% by 8%, 33%, and 29% following the control, statin, and portfolio total fat, 16% SFA), a fruits and vegetables diet (which provided diets, respectively. These results clearly illustrate the effectiveness more fruits and vegetables and fewer snacks and sweets than in the of the portfolio diet over a traditional cholesterol-lowering diet and control diet but otherwise had a similar macronutrient distribution suggest that in controlled settings, the portfolio diet may be as broth), or the DASH diet for 8 weeks. Sodium intake (3000 potent as first-generation statin drugs for lowering LDL-C. mg/day) was similar across all diets, and body weight remained However, this dietary pattern is limited in its effects on HDL-C and constant throughout the intervention.

its effectiveness is reduced. In a 12-month study, subjects were reduced total cholesterol (-9.5 %), LDL-C (-9.1 %), and HDL-C instructed to follow a self-selected low-fat diet (Step II diet) that (-9.2 %), with no change in TG. 21 Smaller reductions in blood incorporated the portfolio of cholesterol-lowering foods (plant pressure were observed in subjects consuming the fruits and sterols, viscous fiber, soy protein, and almonds). ¹⁷ Subjects were vegetables diet (SBP, - 2.8 mm Hg; DBP, - 1.1 mm Hg); total advised to follow a vegetarian diet (without eggs, dairy products, cholesterol, LDL-C, HDL-C, and TG did not change. Stratified by or meat) that included 5 to 10 servings of fruits and vegetables per hypertension status, the DASH diet reduced SBP and DBP by day, with additional plant protein and fiber from dried legumes. If 11.6 mm Hg and - 5.3 mm Hg, respectively, in stage 1 meat or dairy products were consumed, subjects were counseled to hypertensives (SBP > 140 mm Hg, DBP > 90 mm Hg, or both), choose options with reduced SFA and cholesterol. After 12 months, with less dramatic effects in normotensive individuals LDL-C was reduced by 12.8%. This reduction is appreciably less (SBP/DBP, - 3.5/ - 2.2 mm Hg). ²² The most substantial than that achieved in the metabolically controlled study, which is reductions were observed in hypertensive African Americans; probably due to dietary compliance. The authors reported a the DASH diet reduced SBP by - 13.2 mm Hg and DBP by - 6.1 significant correlation between total dietary adherence and change mm Hg. 22 in LDL-C, and by 12 months, the majority of subjects had returned to an omnivorous diet (59 of 66).

evaluated, subjects were most compliant for almonds (79%) and sodium restriction (high, 3200 mg/day; intermediate, 2300 plant sterol-enriched margarine (67%), whereas compliance for mg/day; and low, 1500 mg/day) within a Western or DASH diet. viscous fiber and soy protein was 55% and 51%, respectively. 23 In both the Western and DASH diets, sodium restriction Despite these challenges, 32% of subjects experienced LDL-C progressively lowered blood pressure; however, the effect of reductions of > 20%. The results of this study demonstrate the sodium on blood pressure was more pronounced in subjects difficulty of adhering to a plant-based diet in a free-living setting. following the Western than the DASH dietary pattern (Fig. 16-1). However, for subjects who do achieve this, the portfolio diet is From highest to lowest sodium level, the reduction in SBP/DBP particularly effective at lowering LDL-C. For other individuals, a was 6.7/3.5 mm Hg on the Western diet and 3.0/1.6 mm Hg on focus on the incorporation of individual components (such as the DASH diet. almonds or plant sterol-enriched margarine) may be most efficacious, particularly as plant sterols were identified as the lower blood pressure responses than the Western diet did, with primary LDL-C-lowering dietary component of the portfolio diet. the lowest blood pressure observed in subjects consuming the

CVD risk factors, such as LDL particle size and C-reactive protein consistent across several subgroups, including African (CRP) level. The portfolio diet significantly reduces small LDL-C Americans, hypertensive individuals (SBP > 140 mm Hg or DBP subfractions < 25.5 nm (-0.69 mmol/L); this effect is comparable > 90 mm Hg), older adults (> 45 years), and women, with mean to that achieved by statins and greater than that by a Step II diet reductions of 9.6 to 11.6 mm Hg for SBP and 4.7 to 5.7 mm Hg for alone and translates to a 19% reduction in 13-year risk for CHD. 19 DBP. 24 Interestingly, this study reported that baseline CRP levels predicted

particle size change in response to the portfolio diet. 255 individuals with plasma CRP levels < 3.0 mg/L at baseline showed significant reductions in LDL-C particle < 25.5 nm concentration,

with no change in individuals with CRP levels > 3.0 mg/L. When individuals with high CRP levels (> 3.5 mg/L) were removed from the analysis, CRP level was significantly reduced by statins (16%) and the portfolio diet (24%) but not by the Step II diet (15%). 20 These outcomes illustrate the importance of therapeutic

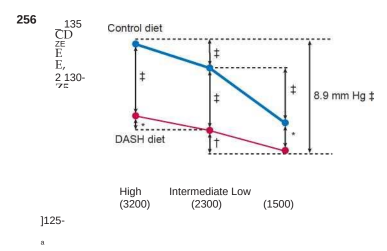
DASH and DASH-Sodium Trials

The DASH trial was a randomized controlled feeding intervention that evaluated the effects of three dietary patterns on blood pressure, lipids, and lipoproteins. 13,21 The DASH dietary pattern,

Compared with a Western diet, the DASH diet lowered SBP When the portfolio diet is implemented in a free-living setting, and DBP by - 5.5 mm Hg and - 3.0 mm Hg, respectively. ¹³ It also

Additional hypotensive benefits can be achieved by following a DASH diet with further reductions in sodium. The DASH-When the individual components of the portfolio diet were Sodium trial compared the hypotensive effects of three levels of

For each level of sodium restriction, the DASH diet elicited DASH diet with the lowest sodium level. The hypotensive effects In addition to effects on LDL-C, the portfolio diet targets other of sodium restriction alone or with the DASH diet were



■ FIGURE 16-1 Change in systolic blood pressure in the DASH-Sodium trial. (Modified from Sacks FM, Svetkey LP Vollmer WM, et al: Effects on blood pressure of reduced dietary sodium and the Dietary Approaches to Stop Hypertension [DASH] diet. N Engl J Med 344:3, 2001. Copyright @2001 Massachusetts Medical Society. All rights reserved.)

W 120-

Sodium intake (mg/day)

* P < .05; f P < .01; J P < .001

Mediterranean-Style Diets

The Mediterranean diet is a whole food-based dietary pattern that has been associated with a reduced incidence of CVD and its associated risk factors. ²⁵ This dietary pattern is representative of the traditions of Crete, Greece, and southern Italy in the 1960s, consumption (one or two 5-ounce glasses/day for men and one 5ounce glass/day for women) was also allowed.

The cardioprotective benefits of the Mediterranean diet may in part be attributed to its nutrient profile, low SFA and high MUFA not differ between groups. This lack of distinction between the two (from olive oil), and high fiber and phytosterol intake (> 400 diets is not surprising given their similar macronutrient mg/day) from plants. Several clinical intervention trials have been composition at 3 months of intervention. undertaken to evaluate the benefits of the Mediterranean diet on PREDIMED Study CVD risk; however, comparisons among studies are challenging as each trial has differed with respect to nutrient profile and the specific foods used to implement the diet.

Lyon Diet Heart Study

The Lyon Diet Heart Study demonstrated that a Mediterranean - and women (60 to 80 years) with diabetes or three or more major type diet, rich in a -linolenic acid, was more effective than a cardiovascular risk factors will be randomized to consume either a modified-fat diet in secondary prevention of coronary events. ²⁶ low-fat diet (total fat < 30%, based on AHA dietary guidelines) or a This study randomized 605 post-myocardial infarction patients to Mediterranean-type diet supplemented with either 30 g/day of consume either a Mediterranean-type diet (30% of total calories mixed nuts (15 g walnuts, 7.5 g hazelnuts, and 7.5 g almonds) or 1 from fat, 8% from SFA, 13% from MUFA, and 5% from PUFA and L/week of olive oil. Participants will be observed for a median 203 mg/day of cholesterol) or a modified-fat diet consistent with a duration of > 5 years and evaluated for primary clinical outcomes Step I diet with a mean follow-up of 46 months. Patients in the (car diovascular death, myocardial infarction, and stroke). Mediterranean type diet intervention group received instructions from the study dietitian and cardiologist to increase their intake of evaluated for changes in lipids and lipoproteins, blood pressure, bread, fish, and root and green vegetables; to consume fruit daily; and glucose after 3 months of intervention. ²⁹ Compared with the to eat less meat (beef, lamb, and pork to be replaced with poultry); low-fat group, the Mediterranean diet with nuts reduced total and to replace butter and cream with margarine supplied by the cholesterol by 6.2 mg/dL and TG by 13 mg/dL. HDL-C was study. The fatty acid composition of the margarine was similar to increased by 2.9 mg/dL and 1.6 mg/dL on the Mediterranean diet that of olive oil, except that it was higher in linoleic acid (16.4% with olive oil and nuts, respectively. Both diets also produced

Control group subjects followed a diet that provided 34% of calories from fat, 12% from SFA, 11% from MUFA, and 6% from PUFA and 312 mg/day of cholesterol. After 48 months, subjects consuming the Mediterranean-type diet (n = 219) had a 50% to 70% lower risk of mortality from heart disease than did subjects consuming a low-fat diet (n = 204). These outcomes occurred despite similar plasma lipids and lipoproteins, blood pressure, body mass index, and smoking status in the two groups, indicating that other risk factors, such as thrombogenesis, are involved in coronary protection.

Medi-RIVAGE Study

The Mediterranean Diet, Cardiovascular Risks and Gene -Polymorphisms (Medi-RIVAGE) study is a 12-month, parallel dietary intervention trial designed to compare the effects of a Mediterranean-type diet and a low-fat diet on CVD risk factors. ²⁷ The low-fat diet was based on AHA guidelines and aimed for a

total fat intake of < 30% of total calories with equal contributions (10% each) from SFA, MUFA, and PUFA. A higher dietary fat intake (35% to 38% of total calories) was recommended for subjects in the Mediterranean-type diet arm, with an emphasis on MUFA (18% to 20% of total calories) from olive oil. SFA and PUFA each contributed 10% of total calories. The low-fat and Mediterraneantype diets were similar in their relative contributions from protein (\sim 15%), carbohydrate (\sim 55% to 60%), and cholesterol (< 300 mg/day). However, the Mediterranean-type diet was higher in fiber (25 g/day compared with 20 g/day in the low-fat group) and allowed two glasses of wine per day for men and one glass for women. Alcohol was to be avoided in the low-fat group.

After 3 months of intervention, both groups (total n = 212) had when the incidence of chronic disease was substantially lower than similarly reduced their total fat, SFA, and cholesterol intake. ²⁸ The that of other countries. Key components of this dietary pattern only macronutrient that differed between the groups was MUFA, include olive oil as the main dietary fat source, abundant plant which was higher in the Mediterranean type diet group (15.6%) foods (fruits, vegetables, grains, cereals, nuts, and seeds), fish and than in the low-fat diet group (13.4%). Significant reductions in shellfish, dairy in low to moderate amounts, poultry, eggs, and total cholesterol were observed in both groups (Mediterraneanlimited amounts of red meat and sweets. Moderate wine type diet, - 7.5%; low-fat diet, - 4.5%), with a trend toward reductions in LDL-C (Mediterranean -type diet, - 11.4%; low-fat diet, - 5.0%). Compared with baseline, both groups showed a decrease in triglycerides, glucose, and insulin, although these did

The Prevention con Dieta Mediterrânea (PREDIMED) study is an ongoing multicenter clinical trial designed to evaluate the effects of a Mediterranean diet on the primary prevention of CVD in 7000 asymptomatic subjects (www.predimed.org). Men (55 to 80 years)

A pilot group of 772 participants (339 men, 433 women) was versus 8.6% kcal) and a -linolenic acid (4.8% versus 0.6% kcal). favorable changes in SBP/DBP (nuts, -7.1/ - 2.6 mm Hg; olive oil, - 5.9/ - 1.6 mm Hg) and glucose concentration (nuts, - 5.4 mg/dL; olive oil, - 7.0 mg/dL) compared with the low-fat diet. Although it appears that a

received simple dietary advice only.

Dietary Patterns That Emphasize Specific Macronutrients

OmniHeart Trial

The OmniHeart trial was a three-period, 6-week crossover, controlled feeding study involving 164 prehypertensive or stage 1 hypertensive subjects that evaluated the cardiovascular benefits of substituting SFA with carbohydrate, protein, or unsaturated fat. 30 Each diet period emphasized the intake of one specific macronutrient – high carbohydrate (58% of total calories), moderate to high protein (25% of total calories, 50% of which were from plant proteins), or high unsaturated fat (31% of total calories, predominantly MUFA) - thereby enabling direct comparison of these macronutrients. Increased unsaturated fat intake caused a 9.2% reduction in TG and a 10.3% reduction in LDL-C but no change in HDL-C (-0.6 %). Protein elicited a similar response, although this was more effective than unsaturated fat in lowering TG (-16.2 %) and LDL-C (-11.0 %) but also caused a modest reduction in HDL-C (-5.2 %).

In comparison, the high-carbohydrate diet was effective only in reducing LDL-C (-9.0 %) and was accompanied by a reduction in HDL-C (-2.8 %). Compared with baseline, all diets decreased SBP (- 8.2, - 9.5, - 9.3 mm Hg, carbohydrate, protein, and unsaturated fat, respectively) and DBP (-4.1, -5.2, -4.8 mm Hg, carbohydrate, protein, and unsaturated fat, respectively); however, on stratification by hypertension status, subgroup analysis showed that in both prehypertensive and stage 1 hypertensive subjects, the moderate- to high-protein and unsaturated-fat diets reduced blood pressure more than the carbohydrate diet did. On the basis of these results, partial replacement of SFA with protein or unsaturated fat appears to be more effective than replacement with carbohydrate in improving lipids and reducing blood pressure.

DIETARY STRATEGIES FOR WEIGHT LOSS

Dietary guidelines issued by the US government and other prominent health organizations, such as the US Department of Agriculture (USDA) ¹¹ and the AHA, ³¹ emphasize the importance of achieving and maintaining a healthy body weight for prevention of chronic disease. However, there continues to be some debate as to the most effective way to achieve weight loss and subsequently to maintain weight loss. In the design of dietary interventions for weight loss, one important aspect that has emerged in recent years is the relative contribution of fat, protein, and carbohydrate. The macronutrient profile of a diet may facilitate dietary adherence, thereby promoting weight loss, and it could promote specific PREMIER Trial changes in blood pressure, lipids, and lipoproteins as the results Recommendations for hypertension management include weight from the OmniHeart trial suggest. Thus far, clinical intervention loss, sodium reduction (< 100 mg/day), increased physical trials have yielded mixed results, and studies are often thwarted by small subject numbers and inadequate follow-up; few studies have evaluated outcomes beyond 1 year, when weight regain is most pronounced. The following section discusses CVD risk factor outcomes from selected weight loss trials. In reviewing these Infarction Early Remodeling) trial evaluated the effectiveness of dietary effects, it is important to acknowledge the role that these management strategies in combination, with or without the additional strategies, such as physical activity and behavioral DASH diet, on blood therapy, may have played in improving weight loss outcomes and preventing weight regain. 32

Mediterranean diet is advantageous, the results of the PRE DIMED The Diabetes Prevention Program (DPP) compared the efficacy of study should be interpreted with caution as both Mediterranean three treatments in preventing type 2 diabetes and metabolic intervention groups received intensive behavioral counseling and syndrome in individuals at high risk on the basis of elevated fasting nutrition education intervention, whereas the low-fat group blood glucose concentration and impaired glucose tolerance. 33,34 Participants (n = 3234) were randomized to receive either standard lifestyle recommendations (written material advising lifestyle changes) plus placebo or metformin (850 mg twice daily) or intensive lifestyle intervention (education curriculum covering diet, exercise, and behavior modification), which aimed to achieve 7% weight loss through a healthy low-calorie, low-fat diet (< 25% of total calories) and 150 min/wk of physical activity.

> At 1 year, participants in the lifestyle intervention group had lost substantially more weight (~ 6.5 kg) than either the placebo (~ 0 kg) or metformin (~ 2.2 kg) groups. Weight loss was a strong predictor of reduced diabetes incidence; each I kilogram of weight lost contributed to a 16% reduction in I risk, 35 and diabetes incidence was 58% and 31% lower in the lifestyle and metformin groups compared with placebo. Intensive lifestyle therapy was also most effective in preventing the development of metabolic syndrome; at 3 years, incidence rates for metabolic syndrome were 38% for the lifestyle group, 47% for metformin, and 53% for placebo. The low-fat diet (as part of the lifestyle intervention) reduced the prevalence of the individual criteria of metabolic syndrome (ie, HDL-C was increased; TG, blood pressure, and glucose levels were reduced).

The Look AHEAD (Action for Health in Diabetes) trial, an ongoing multicenter clinical trial modeled on the DPP, is investigating the long-term effects (11.5-year follow-up) of an intensive lifestyle intervention program in 5145 individuals with type 2 diabetes. ³⁶ This lifestyle intervention program is similar to the DPP in that it combines dietary modification, behavior therapy (based on programs employed in the DPP), and increased physical activity (175 min/wk moderate exercise) to achieve a 10% (minimum 7%) reduction in body weight. However, whereas the DPP focused on fat restriction to decrease calories, the primary method to achieve weight loss in Look AHEAD is through calorie restriction (although fat intake is considered; total fat < 30% of calories, SFA < 10% of calories). In addition to the "toolbox" of adherence strategies used in the DPP, Look AHEAD encourages weight loss medication (orlistat) for patients who do not meet their weight loss goals in the first 6 months. Patients randomized to the intensive lifestyle program are compared with those receiving usual care of diabetes support and education.

After 1 year of intervention, participants in the lifestyle group lost 8.6% of body weight, compared with 0.7% in the diabetes education group. Substantially improvements in glucose control (- 21.5 mg/dL), SBP and DBP (- 6.8/ - 3.0 mm Hg), TG (- 30.3 mg/dL), and HDL-C (+ 3.4 mg/dL) were observed in the lifestyle intervention group. The primary determinants for weight loss success were greater selfreported physical activity, attendance at sessions , and consumption of meal replacements. 37

activity (180 min/wk), moderate alcohol consumption for those who drink alcohol (two drinks/day for men and one drink/day for women), and the DASH diet 6 (see Table 16-3 for approximate reductions in SBP). The PREMIER (Prevention of Myocardial

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were randomized to one of three intervention groups: advice only 12% and 14%, respectively. 41 (control), established lifestyle recommen dations (as described before), or DASH plus established life style recommendations. At 6 Very-Low-Fat Diets months, weight loss was significantly greater in the established (- Multicenter Lifestyle Demonstration Project 4.9 kg) and DASH plus established (-5.8 kg) groups compared The Multicenter Lifestyle Demonstration Project (MLDP) was a with control (-1.1 kg), although both groups experienced weight regain by 18 months (+1.1-1.5 kg). 39

weight men

Similar group treatment effects were observed at 6 months for blood pressure, which was reduced by 10.5/5.5 mm Hg in the established group and 11.1/6.4 mm Hg in the DASH plus established group. However, by 18 months, these changes were not significantly different compared with those observed in the adviceonly group, although this is likely due to the unexpected improvement in several lifestyle factors in this group rather than to to nonfat dairy products and egg whites. Other intervention the limited effectiveness of the interventions.

Fasting insulin, glucose, lipid, and lipoprotein changes have been reported in a secondary analysis that stratified individuals according to the presence of metabolic syndrome. 40 This analysis revealed highly variable effects that differed by treatment group and metabolic syndrome status. The established and DASH plus established interventions compared with the control group lowered total cholesterol in subjects with (- 7.98 and - 5.91 mg/dL, respectively) and without (- 7.41 and - 7.06 mg/dL, respectively) metabolic syndrome. The established intervention reduced TG in both subpopulations (log transformed: with metabolic syndrome, -0.16 mg/dL; without, - 0.10 mg/dL) but lowered LDL-C only in individuals without metabolic syndrome (- 6.89 mg/dL). The DASH plus established group also lowered LDL-C only in individuals without metabolic syndrome (-5.13 mg/dL). Neither intervention influenced HDL-C. Insulin resistance (as determined by homeostasis model assessment) was improved by both the established and DASH plus established interventions, in subjects with and without metabolic syndrome. Despite such variable effects on independent cardiovascular risk factors, these combined Popular Diets behavioral interventions have a profound influence on CHD risk. The majority of popular diets promote extreme carbohydrate or fat

pressure and weight loss. 38 Individuals with untreated, ele vated Compared with advice alone, the established and DASH plus blood pressure (SBP 120 to 159 mm Hg and DBP 80 to 95 mm Hg) established interventions reduced estimated 10-year CHD risk by

comprehensive intervention that examined the combined effects of diet, exercise, stress management, and group support on medical and psychosocial characteristics in 440 patients with coronary artery disease (CAD). 42 The dietary component of MLDP consisted of a low-fat (< 10% of total calories), high-carbohydrate (70% to 75% of total calories), moderate- protein (15% to 20% of calories), predominantly vegetarian diet that emphasized fruits, vegetables, grains, legumes, and soy products. Animal products were limited components included participation in 1 hr/day of stress management, 3 hr/wk of moderate exercise (according to American College of Sports Medicine guidelines for CAD patients), and twice-weekly group support sessions.

At 3 months, subjects had reduced their total fat intake to $\sim 7\%$ of total calories and increased time spent in physical activity (~3 to 4.0 hr/wk) and stress management (~ 5.5 hr/wk). Body weight was reduced by 4 kg in men and 4.6 kg in women. Significant improvements in other cardiovascular risk factors were achieved; blood pressure (men, - 5/ - 5 mm Hg; women, - 6/ - 3 mm Hg), total cholesterol (men, - 18 mg/dL; women, - 14 mg/dL), and LDL-C (men, - 19 mg/dL; women, - 17 mg/dL) were reduced. These behavioral changes and cardiovascular risk factor improvements remained at 12 months. Subjects experienced a substantial reduction in angina (from 42% at baseline to 20% at 1 year for men and 53% to 27% in women), and 20% of individuals with diabetes mellitus reduced their use of glucose-lowering medication. 43

restriction to achieve weight loss. Three recent trials have evaluated the efficacy of popular diets (with significantly different macronutrient compositions) for weight loss and improvements in CVD risk factors. In a 12-month randomized controlled trial involving 160 overweight or obese individuals, Dansinger and colleagues 44 compared the effects of four popular diets: Atkins (very low carbohydrate), Zone (macronutrient balance), Ornish (very low fat), and Weight Watchers (calorie control). The lowcarbohydrate diets (Atkins and Zone) achieved the greatest shortterm (2 month) reductions in TG, glucose concentration, and DBP, whereas reductions in LDL-C and total cholesterol were observed in the higher carbohydrate groups (Ornish and Weight Watchers). These between-group differences were not present at 12 months. Short- and long-term weight loss (2-month range, 3.5 to 3.8 kg; 12month range, 2.1 to 3.3 kg) were similar for all diet groups; the magnitude of weight loss was strongly associated with dietary adherence rather than with diet type.

The A to Z weight loss study ⁴⁵ compared the effects of three popular diets (Ornish, Zone, and Atkins) and a conventional lowfat (< 10% of energy from SFA), high-carbohydrate diet (Lifestyle, Exercise, Attitudes, Relationships, and Nutri tion [LEARN]) on weight loss in 311 premenopausal women.

weight loss (2 months, 4.4 kg; 6 months, 5.6 kg) than did women C. 49 following the Ornish, Zone, and LEARN diets, although only significantly different from Ornish).

Preventing Overweight Using Novel Dietary Strategies (POUNDS LOST) study, a 2-year trial comparing the effects of three primary foods that include unsaturated fats, proteins, or complex macronutrients on weight loss. The study design allowed direct carbohydrates will accentuate the health benefits gained by the comparison of two levels of fat (20% and 40%) and protein (15% reduction in SFA and trans- fat intake. 30 and 25%) intake and a dose response evaluation of carbohydrate (35% to 65% of total calories). All diets were reduced by 750 kcal from baseline and were standardized to include 8% of calories from SFA, at least 20 g/day of dietary fiber, and < 150 mg of cholesterol per 1000 kcal.

Weight loss was similar for participants in each group at 6 months (~ 6 kg, 7% of initial weight) and 2 years (2.9 to 3.6 kg, intention-to-treat analysis). Dietary analysis showed that participants were equally compliant across the four diets; however, in general, participants failed to reach the required calorie reduction and specific macronutrient targets. At 2 years, LDL-C decreased more with the low-fat diets and the highest carbohydrate diet than with the high-fat diets or the lowest carbohydrate diet. The lowest carbohydrate diet increased HDL-C compared with the highest carbohydrate diet. TG, insulin, and blood pressure levels were reduced by all diets (except the highest carbohydrate diet did not lower insulin). These changes are not entirely consistent with the macronutrient effects reported in the OmniHeart study, which may be due to the substantial macronutrient overlap between the diet groups. The results of these three studies collectively suggest that calorie restriction is more important than macronutrient distribution for weight loss; however, modification of macronutrient intake may be instrumental for targeting cardiovascular risk factors such as elevated blood pressure, lipids, and lipoproteins.

KEY FOODS AND NUTRIENTS

The dietary patterns showcased in the previous section have a number of similarities: a focus on weight control; reduced intake of SFA, trans -fat, and cholesterol; and increased intake of fruits, vegetables, whole grains, protein, and reduced-fat dairy. In combination with a heart-healthy dietary pattern, these components have profound effects on blood pressure, lipids, and lipoproteins; yet there is evidence for their independent benefits. The inclusion of other dietary factors, such as increased intake of unsaturated fats, sterols and stanols, soy protein, and fiber, allows greater flexibility for personal preference and additional targeting of specific risk factors. For patients who require a gradual introduction to dietary change, incorporation of one or a few of these strategies may elicit beneficial effects on CVD risk factors.

Key Dietary Components

Cholesterol, Saturated Fatty Acids, and Trans—Fatty Acids

Diets high in SFA and trans fat contribute to elevated levels of total Whole Grains cholesterol in the blood. 48 Sources of SFA are high-fat and processed meats, poultry or game (with skin), high-fat dairy fruit of the grains whose principal components – the starchy products, butter, gravy, and palm and coconut oils.

Trans -fats are commonly found in processed and fast foods, 259 shortening, margarines, fried tortilla chips, commercial or storebought baked goods, salad dressings, candy, and energy bars. Substituted for carbohydrate, SFA increased both LDL-C and

Women following the Atkins diet achieved greater short-term HDL-C, whereas trans-fats increased LDL-C and decreased HDL-

Dietary cholesterol is naturally present in any animal food differences between the Atkins (4.7 kg) and Zone (1.6 kg) diets were source. Dietary cholesterol raises LDL-C; however, its effects are significant at 12 months. Again, for each diet, weight loss was less than those of SFA and trans-fatty acids. 50 Whereas total fat per associated with greater dietary adherence. 46 By 12 months, the se does not affect LDL-C (ie, the guidelines are built on specific fatty Atkins diet produced substantially greater changes in TG (- 29.3 acid recommendations to meet LDL-C goals), it is prudent to mg/dL, significantly different from Zone), HDL-C (+4.9 mg/dL, control total fat intake within current recommended ranges significantly different from Ornish), SBP (- 7.7 mm Hg, (because fat is the most concentrated energy source in the diet) to significantly different from all other diets), and DBP (-4.4 mm Hg, achieve weight loss. Weight loss can lower LDL-C by 5% to 8%. 9 Therefore, reducing the intake of foods that increase LDL-C or Sacks and colleagues 47 recently published outcomes from the decrease HDL-C is an important part of a heart-healthy diet. Furthermore, replacing SFA and trans -fats in the diet with healthy

Fruits and Vegetables

Fruits and vegetables are nutrient-dense and low-calorie 16 foods that are an essential part of a healthy diet. The most abundant nutrients in fruits and vegetables are vitamin C, vitamin E, vitamin A, folate, fiber, and potassium. Many of these are antioxidants and reduce oxidative stress in the body by neutralizing damaging free radicals. Studies using antioxidants in supplement form have not been able to demonstrate their value for reduction of CVD risk 51; interactions between the antioxidants and other nutrients in the whole food may be necessary to have the desired effect. For protection against chronic disease and to maintain good health, the Dietary Guidelines for Americans, 2005 11 recommends consumption of at least 4.5 cups (nine servings) of fruits and vegetables per day for the reference 2000-calorie level (higher or lower amounts can be consumed for other calorie levels).

Fruit and vegetable intake consistently has been associated with decreased risk for CVD in epidemiological studies. In an analysis of more than 125,000 participants from the Nurses' Health Study and the Health Professionals Follow-up Study, persons who ate eight or more servings of fruits and vegetables per day had a 20% reduction in risk of CHD (RR = 0.80; 95% CI, 0.69-0.93) compared with those who ate three or fewer servings per day. 52 With each increase in serving of fruit or vegetable per day, the risk of CHD decreased by 4%; leafy greens and fruits and vegetables high in vitamin C contributed most to this effect. Consistent with these data, the Physicians' Health Study found that men who consumed two or more servings per day of vegetables had a 22% lower risk of CHD than did men who ate less than one serving per day (RR = 0.77; 95% CI, 0.60-0.98). ⁵³ Furthermore, for each additional serving of vegetables per day, CHD risk was reduced by 17% (RR = 0.83; 95% CI, 0.71-0.98). In the Women's Health Study, fruit and vegetable consumption was inversely associated with CVD risk (RR = 0.45 for highest versus lowest quintile of intake; 95% CI, 0.22-0.91). 54 Whereas the epidemiological studies are convincing, there is limited clinical evidence because of the paucity of controlled, nutritional prevention trials varying only in fruit and vegetable intake. 55 Neverthe less, the DASH dietary intervention studies (described earlier) clearly demonstrated the benefit of increasing fruit and vegetable intake for the management of hypertension. 13 In addition, because of their low energy density, fruits and vegetables may play a key role in weight loss and maintenance.

Whole grains are defined as intact, ground, cracked, or flaked

260 endosperm, germ, and bran-are present in the same relative proportions as they exist in the intact grain. 57 If they are processed correctly, wheat, oats, barley, brown and wild rice, corn, rye, and sorghum are whole grains. To be considered a good source of whole grains, foods must be at least 51% whole grain by weight per reference amount commonly consumed and have a whole-grain source as the first ingredient on the food label. Whereas all whole grains have many bioactive components (such as fiber, folate, phenolic compounds, lignan, and sterols), each type of grain has different levels of these components, and therefore they cannot be equated. For example, oats, barley, bulgur, rye, and whole wheat are high in fiber (> 10 g fiber/100 g food), whereas brown rice, wild rice, corn, and sorghum are lower improving CVD risk factors. Specifically, soluble (viscous) section on soluble (viscous) fiber for additional information.

studies have shown whole-grain intake to be protective vitamins. 63.65 against CHD.

CVD, these results may be confounded by lifestyle characteristics. Jensen and colleagues 60 reported that individuals who had higher intakes of whole grains generally Overall, epidemiological studies of dairy intake have shown no had a lower body mass index, were more physically active, protein.

Randomized controlled trials designed to evaluate whether whole grains are protective against CVD generally have shown a beneficial effect, albeit with mixed results. In a recent review of both observational and intervention studies, barley were more effective than whole wheat in improving low dairy consumers. 68 CVD risk factors such as total cholesterol, LDL-C, and blood attributed to the composition of the grain, as oats and barley are higher in fiber, especially insoluble fiber, and have unique servings of whole grains per day is an attainable health goal. or non-fat dairy products within a heart-healthy diet. 13

Reduction of meat intake, especially red meat, is often recommended to decrease intake of SFA and cholesterol. Animal products are the predominant source of SFA and cholesterol in the diet, but processed foods like fast food and snack foods contribute significantly. 62 Animal products are also excellent sources of protein, and increased protein intake, from plant or animal sources, has been linked to lower blood pressure and TG, higher HDL-C, and a healthier body weight. ^{30.63} Protein is low in energy density (4 kcal/g), and therefore replacement of fatty acids with protein may help with satiety and weight loss.

The Nurses' Health Study found that women with a higher protein intake (~ 24% of total energy) were at reduced risk for ischemic heart disease (RR = 0.75; 95% CI, 0.61-0.92) compared with in fiber (<8 g fiber/100 g food). ⁵⁷ Soluble and insoluble fibers women with lower intakes (~14% of total energy). ⁶⁴ However, in have different effects on and mechanisms of action in an analysis of the nutritional profiles of 18- to 30-year old men and women from the Coronary Artery Risk Development in Young fiber has been shown to reduce LDL-C levels, 58 whereas Adults (CARDIA) study, adults who ate meat or poultry less than insoluble fiber may increase short chain fatty acid synthesis, once per week had lower serum TG, total cholesterol, and LDL-C which reduces endogenous cholesterol production. ⁵⁹ See the levels than did those who ate meat more frequently, although the low-meat intake group also had a lower body mass index, increased Similar to fruit and vegetable intake, epidemiological reported physical activity, and a diet higher in fiber and certain

Until recently, there have been no specific recommendations for 16 These associations may be due to the various vitamins (B and intake of protein from animal products with regard to CVD health, E), minerals (calcium, magnesium, potassium, phosphorus, other than to consume reduced-fat products. Protein intake of 10% selenium, manganese, zinc, and iron), phytochemicals (such to 20% of total daily calories from either plant or animal sources is as fiber), phenolic compounds, and phytoestrogens (lignans) widely accepted 63; The Dietary Guidelines For Americans, 2010, found in whole grains. An analysis of the Health Profession however, emphasizes the importance of plant-based dietary as Follow-up Study, which included more than 40,000 men, patterns for CVD risk reduction. 65a For all individuals, the found an inverse association between higher habitual whole challenge is finding good sources of low-fat protein in the diet. grain intake and incidence of CHD (HR = 0.82; 95% CI, 0.70 Meats that are high in SFA are high-fat red meat cuts and processed 0.96); indeed, there was a 6% decrease in CHD risk with every meats, such as hamburger, hotdogs, and bacon, and these should be increase of 20 g in daily whole-grain intake (95% CI, 0%-13%). limited in the diet. Full-fat dairy products are also a rich source of 60 Higher intakes of whole grains (lowest versus highest SFA. Low-SFA animal protein sources are lean meats (beef, ham), quartiles) have been associated with a lower incidence of poultry (trimmed and without skin), fish, cottage cheese, and metabolic syndrome, fasting plasma glucose concentration, reduced-fat milk and dairy products. Plant protein may provide a and body mass index in older adults. 61 Although it appears healthier protein alternative in the diet as it is lower in SFA. that whole-grain intake is associated with a reduced risk for Common sources of plant protein are soy, seeds, nuts, and legumes.

Reduced-Fat Dairy

association with increased risk for CVD, even when full-fat dairy had less hypertension, and ate more fruits and vegetables and products high in SFA are consumed. 66 However, an analysis of the contribution of different fatty acids to CVD risk showed that an increase in the ratio of high-fat to low-fat dairy products increased risk for CVD in the Nurses' Health Study. 67 This suggests either a protective or minimal effect of low-fat dairy products on CVD risk. In a meta-analysis and systematic review of dairy consumption, 11 of 15 intervention studies reported that whole grains were high dairy consumers had a 29% reduced risk for metabolic protective against CVD events. 57 Of note is that oats and syndrome and 14% reduced risk for type 2 diabetes compared with

The main mechanism by which low-fat dairy products reduce pressure. The difference in effect between the whole grains is CVD risk is by lowering blood pressure, although increased insulin sensitivity, decreased inflammation, and weight loss are emerging benefits. Dairy products are an excellent dietary vehicle for protein, phytochemicals. NCEP ATP III recommends 5 to 10 g/day of vitamin A, vitamin D, vitamin B 12, riboflavin, niacin, potassium, soluble (viscous) fiber to lower LDL-C by 3% to 5%. 9 phosphorus, magnesium, and calcium. Increased intake of calcium, Currently, the Dietary Guidelines for Americans, 2005 advise potassium, magnesium, and dairy protein has been shown to that at least half of the recommended grain servings come decrease blood pressure. ^{2.69} According to a meta-analysis of 12 from whole grains and state that "consuming at least 3-ounce" intervention studies, short peptide chains found in milk improve equivalents of whole grains per day can reduce the risk of CVD outcomes by decreasing SBP (- 4.8 mm Hg) and DBP (- 2.2 CHD, may help with weight maintenance, and may lower mm Hg). 70 These nutrients appear to be more effective when they risk for other chronic diseases." 11 Switching from refined to are provided as foods, such as low-fat dairy products, than when whole-grain products is becoming increasingly easier as more they are taken in supplement form. 51 This is a key principle of products are being made with whole grains; thus, three dietary patterns such as the DASH diet, which incorporates low-fat

possible to enjoy the nutritional benefits and to limit SFA intake. any adverse effects. 90 Replacement of high-SFA dairy foods, such as butter and ice cream, achieve a healthy diet and to decrease the risk of CVD. 31

Other Beneficial Dietary Components

Soluble (Viscose) Fiber

Foods rich in water-soluble (viscous) fiber include oats, barley, legumes, some fruits (such as apples and pears), and psyllium seeds. Soluble fibers have been recognized since the 1960s as having lipid-lowering effects, and there is growing evidence to support an Soy Protein association between intake of whole grains and decreased incidence. There has been extensive investigation about the effects of soy of fatal and nonfatal CHD. 60 The cholesterol-reducing effects of whole-grain foods such as oats and barley are associated with the basis of the outcomes of a meta-analysis of 38 trials, 97 the US Food soluble fiber component, beta-glucan.

outcomes, several meta-analyses concluded that regular - cholesterol, may reduce the risk of heart disease." 98 The average consumption of oats can lower cholesterol. 58,71 Brown and col - consumption of soy protein in the studies reviewed in the leagues 58 reported that 3 g/day of soluble fiber decreases total Anderson meta-analysis 97 was 47 g/day, which is almost double cholesterol and LDL-C by - 0.13 mmol/L. Health claims for that recommended in the US health claim. This amount was cholesterol reduction have been approved for both oat fiber 72 and associated with significant reductions in total cholesterol (9.3%), barley fiber. 73 The exact mechanism by which water soluble fibers LDL-C (12.9%), and TG (10.5%). The magnitude of reduction in lower serum LDL-C levels is not known. Water soluble fibers may total cholesterol and LDL-C was associated with baseline levels, interfere with lipid or bile acid metabolism 74 by downregulating such that individuals with higher serum cholesterol levels at genes involved in fatty acid synthesis and transport. 75 Oat bran baseline had the greatest reduction. with beta-glucan has also been shown to increase the exclusion of bile acids. 76 Another suggested mechanism is the inhibition of cholesterol reduction is much less than was initially reported. 99 hepatic cholesterol synthesis by fermentation products. 77 Shortchain fatty acids also may regulate hepatic AMP-activated protein 1999 and found that consumption of isolated soy protein with kinase in the liver, thereby stimulating fatty acid oxidation and isoflavones resulted in a small reduction in LDL-C (3%) but had inhibiting lipogenesis and glucose production. 78 As well as having no effect on HDL-C, TG, lipoprotein(a), or blood pressure. 100 beneficial effects on lowering cholesterol, soluble fiber may reduce Taku and associates 101 found no effect of soy isoflavones CVD risk by acting on glucose regulation and insulin sensitivity, (without concurrent consumption of soy protein) on total body weight, inflammation, endothelial function, and blood cholesterol and LDL-C in menopausal women. Randomized pressure. 79-81

Sterols and Stanols

Several observational studies have evaluated the relationship between plasma phytosterol levels and CVD risk, with two large cohorts reporting a reduced risk of coronary events in individuals with higher plasma sitosterol levels. 82,83 However, the Prospective Cardiovascular Munster (PROCAM) study reported a 1.8-fold increased risk of coronary events in subjects with sitosterol levels in the upper quartile compared with the lower three quartiles. 84 CVD Unsaturated Fats effects may be due to the ability of stands to inhibit cholesterol Both the quantity and quality of dietary fat influence the risk of absorption, thereby reducing plasma total cholesterol and LDL-C. CVD. Intake of unsaturated fats, namely, MUFA and PUFA, is 85 The greatest benefit is observed in individuals with unfavorable associated with a more favorable CVD risk profile as they reduce lipid profiles. 86 In a meta-analysis of supplementation studies, total cholesterol and LDL-C 107 and may improve blood pressure subjects with familial hypercholesterolemia who consumed fat regulation. ¹⁰⁸ Clinical studies have since found that when it is spreads enriched with 2.3 g of phytosterols per day significantly - substituted for SFA in the diet, MUFA reduces total cholesterol reduced their total cholesterol and LDL-C by 7% to 11% and 10% to and LDL-C and relative to carbohydrate increases HDL-C and 15%, respectively, compared with control. 87

significant 5% to 8% LDL-C reduction. 88 A meta-analysis of 41 trials meat, nuts, and avocado. showed that 2 g/day of stanols or sterols reduced LDL-C by 10%, although higher intakes added little to this

effect. 89 This outcome supports the level of plant sterols or 261 stanols (2 g/day) recommended in the NCEP ATP III guide line for lowering of LDL-C. 9 As the typical daily dietary intake of phytosterols in Western cultures ranges from 150 to 400 mg/day, it is necessary to consider dietary supplementation to reach the recommended intake. 89 Phytosterols are well tolerated, can easily

By virtue of reducing the amount of fat in dairy products, it is be incorporated into a range of foods, and are not associated with

Efficacy is similar for sterols and stanols, but food form may with low-fat options, like yogurt and skim milk, provides complete substantially affect LDL-C reduction. 91 Sterols incorporated into fat proteins and essential vitamins and minerals for the maintenance of spreads, mayonnaise and salad dressing, milk, and yogurt appear health. Incorporation of low-fat dairy products into any diet to be more advantageous at lowering LDL-cholesterol than in food increases nutritional quality without markedly increasing calories; products such as croissants and muffins, orange juice, nonfat however, consumption of fat-free dairy foods may limit the amount beverages, cereal bars, and chocolate. The effects of sterols or of fat-soluble vitamins that are absorbed. The DASH diet and TLC stanols on LDL-C lowering are additive with diet or drug advise two or three servings of low-fat dairy products per day to interventions; previous studies reported greater reductions in LDL-C when statin use is combined with foods enriched with plant sterols or stanols 92,93 compared with doubling the dose of statins. 94 Despite such positive effects on cholesterol reduction, sterols I have not been shown to be effective at reducing oxidative stress and endothelial dysfunction, 95 and there are mixed reports in the literature on their ability to reduce low-grade inflammation. 95.96

protein on lipids and lipoproteins during the past decade. 88 On the and Drug Administration approved the following health claim: "25 Although not all individual studies have found positive g of soy protein a day, as part of a diet low in saturated fat and

More recent meta-analyses have concluded that the extent of In 2006, the AHA assessed 22 randomized controlled trials since controlled trials conducted since these reviews have reported similar, modest reductions in LDL-C or total cholesterol. 102,103 The variable effect of soy isoflavones on cholesterol reduction may be due to the different doses of isoflavones used 104 or may be related to the ability of individuals to convert the isoflavone daidzein into equol through bacterial fermentation in the large intestine. 105 However, evidence to support the latter is not consistent. 106

reduces TG. 30,107 MUFAs are mainly found in vegetable oils such At least 1 g/day of phytosterols is necessary to obtain a as olive, rapeseed (canola), and peanut oils as well as in poultry,

found in vegetable oils and nuts, and omega-3 (n-3) fatty acids, found in fatty fish, such as salmon and anchovies, and in walnuts and flax. Three of the main dietary sources of unsaturated fat and their effects on CVD risk are discussed in greater detail.

Fish (and Fish Oil). Numerous epidemiological and clinical studies have demonstrated associations between increased intake of marine-derived n-3 fatty acids from fish or fish oil and reduced incidence of CVD. 109 In the US Health Professionals Study, regular fish consumption was associated with a significantly lower risk of total CVD in men. 110 It also is associated with reduced progression of coronary artery atherosclerosis in men and women with CAD, 111,112 reduced risk of CAD and total mortality in diabetic women, 113 reduced risk of thrombotic infarction, 114 and reduced risk of cardiac arrhythmias resulting in sudden cardiac death. 115 The n-3 fatty acids in fish are the key nutrients thought to be responsible for the benefits described, although it is plausible that the interactions between these fats and other nutrients, including trace elements, vitamins, and amino acids, may also be important to reduce CVD risk. 116 Marine-derived n-3 fatty acids elicit cardioprotective benefits through mechanisms such as antiarrhythmic effects, TG 16and blood pressure lowering, reduced platelet function and aggregation, improved vascular function, and decreased inflammation. Consequently, regular consumption of marinederived n-3 fatty acids is recommended for healthy persons and Data from Kris-Etherton PM, Harris WS, Appel LJ: Fish consumption, fish oil, omega-3 fatty those with CHD.

Recommendations to eat fish and other seafood are included in most national dietary guidelines because of their beneficial health effects. Seafood is an important dietary source of marine-derived n- from circulating very-low-density lipoprotein (VLDL) particles by 3 fatty acids, protein, vitamins D and E, and iodine. Seafood is low increasing lipoprotein lipase activity. Some studies have shown a in SFA. Because of this unique nutrient profile, regular fish small increase in HDL-C (11% to 14%) with high doses of marineconsumption is recommended as part of a healthy diet. The derived n-3 fatty acids (4 g/day). 122 American Dietetic Association and Dietitians of Canada recommend two servings of fish per week, preferably fatty fish. 117 macronutrient and micronutrient profiles that are cardioprotective. The AHA also recommends two servings of oily fish per week 123,124 They have a healthy fatty acid profile, containing high levels (equivalent to 400 to 500 mg of marine-derived n-3 fatty acids per of the 18-carbon MUFA oleic acid and low amounts of SFA, and day) for optimal health. 31,118 However, fish intake in Western - some varieties (eg, Brazil nuts, pine nuts, and walnuts) are cultures is typically very low (about one fish meal per week) and important sources of n-6 and n-3 fatty acids. 125 Nuts are rich in plant frequently comes from sources that are low in marine-derived n-3 protein, fiber, and micronutrients (such as potassium, calcium, and fatty acids (eg, shrimp, cod, and other white fish). 119

16-4. Farmed fish provides as much EPA and DHA as wild fish do, turn elicit potential antiatherogenic effects. if not more. As the fatty acid profiles of diets for farmed fish can be closely monitored, their lipid is less affected by seasonal variations associated with decreased CVD risk in several large, prospective and location of catch. Moreover, the EPA and DHA content of cohort studies including the Adventist Health Study, 127 the Iowa farmed salmon and trout has been reported to be ~ 15% higher than Women's Health Study, 128 the Nurses' Health Study, 129 and the that of wild catch. ¹¹⁸ The nutritional content of farmed fish has been Physicians' Health Study. ¹³⁰ Collectively, the evidence suggests that found to be at least as beneficial as that of wild fish in terms of the risk of CHD is 37% lower for people consuming nuts more than prevention of CVD, with additional benefits arising from greater four times per week compared with those who never or rarely control over pollutants (including heavy metals and polychlorinated biphenyls). 120

is low, particularly for individuals with a history of CHD or TG clearance

PUFAs are composed of two classes: omega-6 (n-6) fatty acids, TABLE 16-4 Amount of Marine-Derived n-3 Fatty Acids (EPA and DHA) in Select **Species of Fish**

Serving, Edible Portion)

1.81

1.71

0.98

0.84

EPA and DHA (g/3-ounce Fish

Pacific

Atlantic

Trout, rainbow

Farmed Wilde

hunder			
light, canned in water, drained	0.26 0.73 0.24-1.28		
Vhite, canned in water, drained			
Fresh			
Sardines	0.98-1.70		
Salmon			
Chum	0.68		
Sockeye	0.68		
Pink	1.09		
Chinook	1.48		
Atlantic, farmed	1.09-1.83		
Atlantic, wild	0.9-1.56		
Mackerel	0.34-1.57		

acids, and cardiovascular disease. Circulation 106:2747, 2002.

Nuts. Nuts are energy- and nutrient-dense foods that have magnesium). Tree nuts and peanuts also contain numerous Oily fish rich in marine-derived n-3 fatty acids include tuna, phytochemicals, including phytosterols, tocopherols, antioxidant salmon, mackerel, sardines, herring, and trout. The amount of vitamins, and flavonoids (eg, resveratrol in peanuts). 125,126 These eicosapentaenoic acid (EPA) and docosahexaenoic acid (DHA) is constituents may reduce oxidative stress and counteract any shown for a variety of fish species (both fresh and farmed) in Table potential pro-oxidative effects of the unsaturated fatty acids and in

Frequent consumption of nuts or peanut butter has been consume nuts, with an average reduction of 8.3% for each weekly serving of nuts. ¹³¹ However, the evidence supporting a reduced risk Fish oil supplementation is advised when dietary intake of fish of heart failure with regular nut consumption is less strong. 132

A range of cardiometabolic health benefits are likely to hypertriglyceridemia (TG levels > 500 mg/dL). The AHA contribute to these epidemiological observations. 124 Both tree nuts recommends 1 g/day or 2 to 4 g/day of marine-derived n-3 fatty and peanuts have lipid-lowering properties. Clinical studies have acids for individuals with CHD or hypertriglyceridemia , demonstrated that when nuts are included as part of a healthy diet respectively, under the supervision of a physician. On the basis of that is low in SFA and cholesterol, they elicit favorable effects on data from a meta-analysis of 72 placebo-controlled trials, this level lipids and lipoproteins compared with a control diet (typically of supplementation (3 to 4 g/day) will reduce TG levels by 25% to either a low-fat diet or an average American/Western diet). The 35%, with more substantial reductions in hypertriglyceridemic primary outcome is a reduction in plasma LDL-C and TG, along individuals. 121 The mechanism by which marine-derived n-3 fatty with an increase in HDL-C. 133 Nuts have also been found to reduce acids lower TG levels is unclear. There is evidence that they increase oxidation and inflammation and to improve glucose regulation and

Olive Oil. Epidemiological and clinical studies have reported protection. that the traditional Mediterranean-style diet is associated with significantly lower mortality from CAD. ²⁶ Although it is difficult to being investigated. Resveratrol is a natural polyphenol present in isolate individual dietary factors, cumulative evidence suggests red wine that has purported atheroprotective properties. 154 There that olive oil, which is the primary source of fat used by is evidence that resveratrol can act through several mechanisms, Mediterranean populations, may play a key role in the observed including inhibition of platelet aggregation 155 and smooth muscle cardiovascular benefit. 138 Olive oil is rich in oleic acid (typically cell proliferation, 156 reduction of LDL oxidation, and enhanced about 75% of total fatty acids), and virgin (unrefined) olive oil cholesterol efflux. 157 Several studies have reported a decrease in the contains a significant amount of antioxidants (a -tocopherol) and susceptibility of LDL I particles to oxidation in healthy subjects after phytochemicals.

development of atherosclerosis. The unsaturated fatty acid profile inconsistency of these results may be related to variations in the 16 of olive oil lowers LDL-C and increases HDL-C. It also contains polyphenol concentration of the red wine used in the different polyphenols that reduce oxidative stress as they are able to studies. It is unclear whether polyphenols from red wine can scavenge free radicals and protect LDL from oxidation. Results of provide additional cardiovascular benefits beyond that of ethanol the EUROLIVE study ¹³⁹ have provided evidence for the protective alone. role of phenolic compounds from olive oil on in vivo LDL oxidation in humans. Regular olive oil consumption has also been shown to be protective against hypertension in some epidemiological studies, including the Seguimiento Universidad de Navarra (SUN) study 140 FOOD-BASED DIETARY RECOMMENDATIONS and the Greek European Prospective Investigation into Cancer and Nutrition (EPIC) study. 141

Omega-6/Omega-3 Ratio. Both n-3 and n-6 fatty acids are essential PUFAs that are necessary for proper growth and development. The n-3 fatty acids in cell membranes produce antiinflammatory, pro-resolving eicosanoids, whereas n-6 fatty acids are the precursors to proinflammatory prostaglan dins. 142 Eicosanoids and prostaglandins affect a number of pathways associated with CVD progression, such as thrombosis, inflammation, and vasoconstriction. Thus, it is hypothesized that a balanced ratio of n-6/n-3 fatty acids will attenuate inflammatory and vasoconstrictive pathways, resulting in reduced CVD risk. MyPyramid was developed by the USDA to incorporate between n-6 fatty acid intake and inflammatory markers. 143,144 Moreover, in a study conducted by Ferrucci and coworkers, 145 certain proinflammatory markers were decreased and an antilevels of EPA and DHA.

to 10% of total daily calories be provided by n-6 fatty acids, and ranges in the number of servings from each group (to account for individuals who consume less may have a higher risk for CVD events. 147 Thus, a higher PUFA diet that emphasizes both n-3 and n-6 fatty acids is the current recommendation to reduce CVD risk.

Red Wine and Alcohol

Epidemiological studies have indicated that moderate red wine consumption is associated with reduced rates of cardiovascular morbidity and mortality. 148 There has been 263 some debate as to whether the protective effects of red wine result from the alcohol content or the antioxidant properties of the flavonoids found at high concentrations in some red wines. 149 The protective effect of alcohol may partly be attributed to alcohol-associated increases in HDL-C

endothelial function, ¹³⁴⁻¹³⁶ and they may be protective against the by stimulation of reverse cholesterol transport pathways ¹⁵⁰ and development of hypertension. 137 However, this area of research is reduced platelet aggregation. 148 Although the concentration of less advanced than that for lipids and lipoproteins , and antioxidant polyphenols is higher in red than in white wine, some consequently the effects of tree nuts and peanuts on oxidative studies have found that both types of wine 151 as well as other stress, inflammation, and blood pressure are generally inconsistent. alcoholic beverages 152,153 provide a similar extent of cardio vascular

The role of resveratrol (*trans* -3,4 ' ,5-trihydroxystilbene) is also the daily consumption of 375 to 400 mL of red wine for 2 weeks, 158 There are multiple mechanisms by which olive oil may affect the although this was not reported in another study. 159 The -

Food guides translate recommendations on nutrient intake into food-based guidance. Such information is presented in a framework that promotes the selection of a variety of foods that together provide a nutritionally adequate diet. These dietary principles help people to maintain and improve their health and to reduce their risk for major chronic diseases. Graphic presentations, such as the pyramid, are often used to convey important guidelines about food intake to help consumers implement dietary recommendations.

MyPyramid

However, a number of studies have reported no relationship recommendations from the Dietary Guidelines for Americans, 2005. Wo The MyPyramid symbol conveys a personalized approach to diet and physical activity for healthy individuals and promotes several key messages: variety, proportion, and moderation (Fig. 16-2). The inflammatory marker was elevated with increasing serum levels of colored bands (orange, green, red, blue, purple) represent the five arachidonic acid (a metabolite of linoleic acid) as well as serum food groups of the pyramid (grains, vegetables, fruits, milk, meat and beans) and oils (yellow) and suggest that a variety of foods be As noted by Harris, 146 the concept of balancing the n-6/n-3 fatty consumed from each of these groups. The varying width of each acids ratio in the diet is conceptually flawed. It over looks the band serves as a guide to the proportion of food that should be absolute levels and chain-length differences within each fatty acid consumed from each band (actual serving size can be determined class. For example, an n-6/n-3 ratio of 5 can be achieved by vastly by visiting www.MyPyramid.gov). For example, the greater width of different amounts of linoleic acid and n-3 PUFAs, the latter of which the orange grain band indicates that people should eat more foods could include widely differing amounts of a -linolenic acid, EPA, from this group than from the purple meat and beans group. and DHA. Currently, it is estimated that the ratio of n-6/n-3 fatty Moderation is represented by the narrowing of each food group acids in the Western diet is 15 to 20: 1. All current recommendations towards the top of the pyramid. Foods that are lower in total or solid advocate that n-3 intake be increased. Thus, the n-6/n-3 ratio as a fats and added sugars form the base of the pyramid and should be measure of diet quality is flawed for a number of reasons. The best chosen more often. The nar rower area represents foods that are advice is to consume adequate amounts of both n-6 and n-3 fatty high in solid fats and added sugar. People who are more active can acids, thereby focusing on the total amounts of both n-3 and n-6 consume more of these foods. Other features of MyPyramid fatty acids. The AHA supports the recommendation that at least 5% designed to assist consumers in choosing a healthier diet include

DASH Pyramid

The DASH diet is low in total fat, SFA, and cholesterol and rich in fruits, vegetables, and reduced-fat dairy products. It also includes whole grains, nuts, fish, and poultry and there fore promotes foods that are rich in blood pressure-lowering



FIGURE 16-2 The USDA Food Guide Pyramid.

nutrients, such as potassium, magnesium, calcium, and fiber. It is reduced in sodium, red meat, sweets, added sugars, and fats. The DASH diet has been modified by the Center for Science in the Public Interest's Nutrition Action Healthlet ter 161 to fit the Food Guide Pyramid model first published by the USDA in 1992. The DASH pyramid has a base of fruits and vegetables (8 to 10 servings/day) and grains (preferably whole grains, 7 to 8 servings/day) at level 2. Level 3 is shared by low-fat dairy (2 to 3 servings/day) and seafood, poultry, and lean meat (0 to 2 servings/day). Vegetables, nuts, and seeds (1 serving/day) and oils (for healthy fats; 2 to 3 servings/day) make up level 4. The final level is composed of sweets (no more than 5 servings/week). Lower salt foods should be chosen from all categories. The DASH pyramid developed by the Center for Science in the Public Interest also provides examples of servings for each food group, which may be a useful resource for meal planning.

Mediterranean Pyramid

The Mediterranean dietary pattern emphasizes vegetables, fruits, grains, nuts and seeds, unsaturated fats (primarily from olive oil), fish, and shellfish. Poultry, eggs, and dairy are eaten regularly but

> amounts. Intake of red meat and sweets is limited. Use of herbs and spices encouraged they add flavor to foods and reduce the need to add salt or fat during cooking. food choices are reflected in the Diet Pyramid (Fig. 16-3). 162 Regular, moderate consumption (one or two 5-ounce glasses/day men

in low to moderate

as

These

Mediterranean

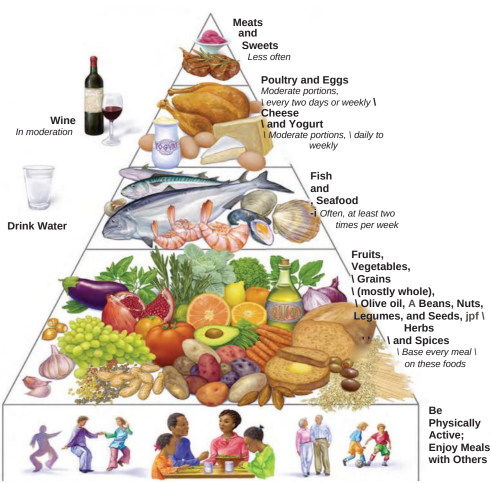


FIGURE 16-3 The Mediterranean Diet Pyramid. (© Oldways Preservation and Exchange Trust. 162)

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and one 5-ounce glass/day for women), participation in daily necessarily translate to reduced risk for stroke, myocardial physical activity, adequate water intake, and enjoying meals in the infarction, or cardiovascular mortality. The VISP (Vitamin company of others are also important components. The visual Intervention for Stroke Prevention) trial randomized 3680 adults display of the Mediterranean Diet Pyramid assists consumers in with nondisabling cerebral infarc tion to receive either high-dose determining portion size and frequency of consumption of various (2.5 mg folic acid, 400 | ig B₁₂, and 25 mg B₆) or low-dose (20 | ig foods; foods in the bottom section of the pyramid may be eaten in folic acid, 6 | ig B 12, and 200 | ig B 6) B vitamins. 171 Despite a larger amounts and more often, whereas the portion size and reduction in homocysteine after 2 years of follow-up, there was no frequency of consumption decline in the upper section of the effect on vascular outcomes. pyramid.

SUPPLEMENTS

Dietary supplement use in the United States is increasing. 163 In the NHANES 1999-2000 survey, 52% of adults reported taking a dietary significant effect on recurrent I CVD in any of the treatment groups. supplement in the last month, the most prominent of which were multivitamin and multimineral supplements (35% of individuals). ¹⁶⁴ However, there are limited data supporting the use of nutritional supplements for CVD risk reduction, and some supplements may even have adverse effects. 51 Several prominent organizations have determined that the evidence for the prevention of CVD by vitamin on overall cardiovascular mortality or combined cardiovascular E, selenium, and multivitamins (containing folic acid) is risk. 173 Other studies also have reported no effect of folic acid (0.8 inconclusive, 165 and the AHA also does not recommend antioxidant mg), B 12 (0.4 mg), or B 6 supplementation (40 mg) on cardiovascular supplementation for CVD risk reduction. 166 The following section events or total mortality in patients with CAD. 174 Homocysteine briefly discusses the evidence for use of select dietary supplements lowering (18.5% more than placebo) with a combination to improve CVD risk factors and to reduce CVD mortality and morbidity.

Niacin

Niacin is a broad-spectrum lipid-regulating drug that has wellestablished effects on CVD risk. In pharmacological doses of 2 to 4 Antioxidants from dietary sources, such as vitamin E, vitamin C, g/day, niacin reduces total cholesterol (20%), TG (25% to 50%), and carotenoids, have been studied for their potential to protect VLDL (45%), LDL-C (5% to 25%), and lipoprotein(a) (40%) and against oxidative stress. Prospective cohort studies have shown increases HDL-C (25% to 50%). 167,168 Niacin, alone or in promising inverse associations between intake of antioxidants, combination with other lipid-lowering agents, significantly reduces particularly carotenoids and vitamin E, and risk for CVD. 176 total mortality and coronary events, retards progression, and However, findings from clinical intervention studies have induces regression of atherosclerosis. ¹⁶⁷⁻¹⁶⁹ However, outcomes reported null and even adverse effects on CVD outcomes from from the HDL-Atherosclerosis Treatment Study (HATS) suggest intake of large doses of single and combination antioxidant that antioxidant vitamins (800 IU vitamin E, 1000 mg vitamin C, 25 supplements. 166,177-179 A recent meta-analysis of 68 randomized mg beta carotene, and 100 µg selenium) may blunt any beneficial trials with 232,606 participants reported that antioxidant effects of simvastatin and niacin treatment on lipids and stenosis supplements taken daily or on alternate days (mean duration, regression. 169 Niacin (1 to 2 g/day) is currently recommended for 2.7 years) had no significant effect on mortality. 177 However, the treatment of elevated TG and LDL-C levels and low HDL-C. 9 when only low-bias risk trials (those with high methodological Despite such broad beneficial effects on lipids and atherosclerosis, quality) were included in the analysis, beta carotene (mean dose, the clinical use of niacin is limited in some patients because of 17.8 mg), vitamin A (mean dose, 20,219 IU), and vitamin E (mean adverse effects, such as flushing and hepatotoxicity. Over-the- dose, 569 IU/day), singly or combined, significantly increased counter formulations that promote "no flushing" vary widely in mortality. There was no effect of vitamin C (mean, 488 mg) or quality and do not have the same benefits as prescription niacin. For selenium (mean, 99 µg) on cardiovascular mortality. These this reason and because of the risks associated with niacin outcomes are supported by more extensive analysis as part of a supplementation beyond standard vitamin doses, it is advised that Cochrane Review of antioxidant supplementation and mortality only prescription niacin be considered and administered under prevention. 178 The AHA advocates the consumption of medical supervision.

B Vitamins (Folate, B 6, B 12)

Elevated levels of homocysteine are associated with an increased risk for ischemic heart disease, stroke, peripheral vascular disease, and thrombosis, possibly because of increased oxidative stress, Vitamin D is a fat-soluble vitamin that is produced by skin thrombosis, and vascular dysfunction. Several factors, including an exposed to ultraviolet B rays or is found in foods like salmon and inadequate dietary intake of B vitamins (folic acid, B₁₂, and B₆), fortified milk. The main function of vitamin D is to maintain have been associated with increased homocysteine concentrations, appropriate calcium and phosphorus levels in the blood and supple mentation with B vitamins may therefore provide a means to reduce CVD risk.

A meta-analysis of 25 randomized controlled trials reported that 0.2 to 5.0 mg/day of folic acid reduced homocysteine by

13% to 25%. 170 The addition of vitamin B $_{12}$ (mean dose, **265** 0.4 mg/day) produced a further 7% reduction, but vitamin B 6 had no effect. However, several long-term clinical trials demonstrate that homocysteine lowering with B vitamin supplementation does not

The NORVIT (Norwegian Vitamin) trial evaluated clinical outcomes during 3.5 years in 3749 patients with a recent myocardial infarction. 172 Patients were randomized to one of four groups and received daily folic acid (0.8 mg) and B vita mins (40 mg B 6 and 0.4 mg B $_{12}$), folic acid (0.8 mg/day) and B $_{12}$ (0.4 mg), B $_{6}$ (40 mg), or placebo. Despite a reduction in homocysteine, there was no A trend towards an increased risk was observed in the combined folic acid and B vitamin group (B 6 and B 12). The Heart Outcomes Prevention Evaluation (HOPE) 2 study reported that daily supplementation with folic acid (2.5 mg), B $_{6}$ (50 mg), and B $_{12}$ (1 mg) in patients with diabetes or vascular disease for 5 years had no effect supplement of folic acid (2.5 mg), B₆ (50 mg), and B₁₂ (1 mg) also failed to reduce cardiovascular events in high-risk women. 175

Antioxidants (Vitamin E, Vitamin C, Carotenoids)

antioxidant-rich foods such as fruits, vegetables, whole grains, and nuts rather than antioxidant supplements for CVD risk reduction. 31

Vitamin D

through the regulation of calcium absorption from the diet, resorption from bones, and reabsorption in the kidneys. Recent evidence suggests that vitamin D also plays a role in neuromuscular function, immune function, blood pressure, and inflammation as well. Low levels of vitamin D in the blood are associated with increased risk for cancer, multiple sclerosis, and Parkinson's

Thus far, clinical trials have not been able to prove causa tion, although there is plausible biological and robust epidemiological evidence to support the role of vitamin D in the prevention of CVD. A large cross-sectional study of patients who were referred for coronary angiography showed that those with vitamin D deficiency (< 25 nmol/L) compared with those with optimal vitamin D levels (> 75 nmol/L) had a hazard ratio of 2.84 (95% CI, 1.20-6.74) and 5.05 (95% CI, 2.13-11.97) for death by heart failure and sudden cardiac death, respectively. 181 In a prospective cohort study (Framingham Offspring cohort) of 1739 adults with no history of CVD, low levels of serum vitamin D (< 15 ng/dL) were associated with an increased risk for cardiovascular events (HR, 1.62; 95% CI, 1.11-2.36) compared with those with normal vitamin D levels (> 15 ng/dL). 182 This relationship was present in individuals with 16 hypertension (2.13; 95% CI, 1.30-3.48) but not in those without hypertension (95% CI, 0.55-1.96). Overall, multiple meta-analyses show a strong link between low vitamin D status and increased CVD risk ^{183,184}; however, the intervention studies designed to test the effect of vitamin D supplementation on CVD outcomes are currently being conducted.

The sun provides much of the vitamin D needed by the body, and 5 to 10 minutes of exposure of bare face, arms, and hands per day should be sufficient to obtain the recommended dietary allowance for vitamin D (200 IU, for those younger than 50 years). ¹⁸⁵ However, this conversion depends on a multitude of factors including skin color, time of year, location, and genetics. Currently, a committee from the Food and Science Board of the Institute of Medicine is reviewing the recent literature with regard to vitamin D recommendations and may increase the dietary reference intake in the coming year. Some experts now recommend 800 to 1000 IU/day of vitamin D 3 from supplements or diet for adults and children to avoid vitamin D deficiency when sunlight exposure is not sufficient. ¹⁸⁶ Because the majority of vitamin D is from the sun measures of serum vitamin D levels (25-hydroxyvitamin D) are important for diagnosis of insufficient or inadequate vitamin D status. Unfortunately, there has been difficulty in standardizing measurement methodology. This is problematic when trying to correlate vitamin D status with health outcomes, and additional research is needed to prove causation between vitamin D status and

AMERICAN HEART ASSOCIATION DIETARY GUIDELINES REVISION 2006

It is evident that healthy diet and lifestyle behaviors markedly decrease the risk of CVD. Consequently, the AHA historically has to heart health. been at the forefront of issuing diet and lifestyle recommendations for heart health. The most recent diet and lifestyle Lifestyle Recommendations are presented in Table 16-5. Specific recommendations were issued in 2006. 31 The 2006 goals are food-based recommendations have been made for the DASH presented in Box 16-1 and target major risk factors for CVD dietary pattern and the TLC diet that promote the AHA Diet and including overweight and obesity, abnormal lipids and Lifestyle Recommendations 2006. These diets are comparable and lipoproteins, high blood pressure, high blood glucose levels, recommend a dietary pattern that is high in vegetables and fruits, sedentary behavior, and exposure to tobacco products.

presented in Box 16-2 and target the major CVD risk factors. vegetable oils. Achievement of a healthy body weight will reduce the risk of

Consume an overall healthy diet.

Aim for a healthy body weight.

Aim for recommended levels of LDL-C, HDL-C, and triglycerides.

Aim for a normal blood pressure.

Aim for a normal blood glucose level.

Be physically active.

Avoid use of and exposure to tobacco products.

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Balance calorie intake and physical activity to achieve or maintain a healthy body

Consume a diet rich in vegetables and fruits.

Choose whole-grain, high-fiber foods.

Consume fish, especially oily fish, at least twice a week.

Limit your intake of saturated fat to <7% of energy, trans-fat to

<1% of energy, and cholesterol to <300 mg per day by: choosing lean meats and vegetable alternatives; selecting fat-free (skim), 1%-fat, and low-fat dairy products; and

minimizing intake of partially hydrogenated fats.

Minimize your intake of beverages and foods with added sugars.

Choose and prepare foods with little or no salt.

If you consume alcohol, do so in moderation.

When you eat food that is prepared outside the home, follow the AHA Diet and Lifestyle Recommendations.

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> CVD and facilitate achievement of a recommended lipid and lipoprotein profile, blood pressure, and glucose level. Limiting SFA, trans -fatty acids, and cholesterol will also help achieve a recommended lipid and lipoprotein profile. Limiting sodium (< 1500 mg/day) will help achieve a normal blood pressure. Fruit and vegetable, whole-grain, and fatty fish recommendations will decrease CVD risk by providing a mineral-rich, fiber-rich diet high in long chain n-3 fatty acids, all of which have been shown to be cardioprotective. Minimizing added sugar will help with calorie control as well. Moderating alcohol consumption can be of benefit

Food-based recommendations consistent with the 2006 Diet and whole grains, fat-free or low-fat dairy products, lean meats and The AHA 2006 Diet and Lifestyle Recommendations are poultry, fish (preferably oily), nuts, seeds, legumes, and liquid

TABLE 16—5 Exam	ples of Dietary Patter	rns That Are Consistent with Americ	an Heart Association Dietary Guidelines*	
Food Group	DASH	Eating Pattern	USDA Dietary Guidelines	Serving Size
Vegetables	4-5 servings/day	5 servings/day	5 servings/day	1 c raw leafy vegetables, 12 c cut-up raw or cooked vegetables, 12 c vegetable juice
Fruits	4-5 servings/day	4 servings/day	4 servings/day	1 medium fruit; / ₄ c dried fruit; 12 c fresh, frozen, or canned fruit; 12 c fruit juice
Grains	6-8 servings/day	7 servings/day	7 servings/day	1 slice bread, 1 oz dry cereal, 12 c cooked rice or pasta
Fat-free or low-fat milk products	2-3 servings/day	2-3 servings/day	3 servings/day	1 c milk, 1 c yogurt, % oz cheese
Lean meats, poultry, fish	<6 oz/day	<5 oz/day	5.5 oz/day	1 oz cooked
Nuts, seeds, legumes	4-5 servings/day	Counted in vegetable servings	Counted in lean meats, poultry and fish, and vegetable servings*	13 c, 2 tbsp peanut butter or seeds, 12 c dry beans or peas
Fats and oils	2-3 servings/day	Amount depends on daily calorie level	Liquid oils: amount depends on daily calorie level Solid fats: counted in discretionary calorie allowance (—10% total kcal)	tsp soft margarine, 1 tbsp mayonnaise, tsp vegetable oil, 2 tbsp salad dressing
Sweets and added sugars	<5 servings/day	No recommendation	Counted in discretionary calorie allowance (— 10% total kcal)	1 tbsp sugar, jelly, or jam; 12 c sorbet; 1 c lemonade

^{*}For daily calorie intake of 2000 kcal.

Modified from Lichtenstein AH, Appel LJ, Brands M, et al: Diet and lifestyle recommendations revision 2006: a scientific statement from the American Heart Association Nutrition Committee. Circulation 114:82, 2006. @2006, American Heart Association, Inc.

AMERICAN HEART ASSOCIATION STRATEGIC IMPACT GOALS THROUGH 2020

The AHA 2020 Impact Goals are "to improve the cardiovascular health of all Americans by 20% while reducing deaths from cardiovascular diseases and stroke by 20%." 12 To meet this goal, the AHA recommends the consumption of an overall healthy dietary pattern that is consistent with a DASH-type eating plan. Key components include but are not limited to the following:

- Fruits and vegetables: > 4.5 cups per day
- Fish: two or more 3.5-ounce servings per week (preferably oily
- Fiber-rich whole grains (> 1.1 g of fiber per 10 g of carbohydrate): three or more 1-ounce equivalent servings per day
- Sodium: < 1500 mg/day
- Sugar-sweetened beverages: < 450 kcal (36 ounces) per week. Secondary dietary goals that support cardiovascular health and are consistent with a DASH-like diet include the following:
- Nuts, legumes, and seeds: > 4 servings per week
- Processed meats: none or < 2 servings per week
- Saturated fat: < 7% of total energy intake

SIMPLE STRATEGIES TO IMPLEMENT AND TO MAINTAIN HEALTHY **DIETARY CHANGES**

Behavior change is central to the achievement of diet and life style recommendations for heart health. Much research is being done to identify effective behavior change strategies, and there are some strategies that seem to work. It all starts with knowing if a patient

is "ready" to make changes in a sustained way. If not, then work is needed to "get" the individual to a stage of change state. The Stages of Change model 187 is an integrative model of intentional behavior change. It is used to describe how people can achieve a desirable behavior or modify a problem behavior. In attempting to change a specific behavior, a person typically cycles through a series of five stages: pre-contemplation, contemplation, preparation, action, and maintenance. Identification of what stage an individual is in will assist in determining what techniques will be most successful in moving the individual to the next stage of change. Motivational interviewing is widely used in practice to evaluate the stage of change and to enhance intrinsic motivation to change by exploring and resolving ambivalence. 188 In addition, motivational interviewing is effective at identifying reluctance to change and helps the individual "see" that needed change can be achieved.

Presuming the individual is ready to change, important steps to follow include goal setting (ensuring that goals are realistic), selfmonitoring strategies (such as keeping food records, body weight measurements), and ensuring that social support is available. These strategies promote self-efficacy (ie, helping patients see that they can achieve short-term goals and pursue longer-term goals). Inherent to modifying behavior is that change strategies need to be individualized and monitored closely for behavior change to be sustained. We encourage readers to refer to a recent scientific statement published by the American Heart Association, which provides an excellent review of interventions to promote physical activity and dietary lifestyle changes for the reduction of CVD risk factors. 189

NATIONAL AND COMMUNITY **INITIATIVES AND RESOURCES**

Many resources are available to the public to implement current diet and lifestyle recommendations for heart health. The government is actively promoting educational programs that promote heart health. Likewise, professional

268 organizations have many programs available for different target groups. In addition, the private sector is committed to improving the health of Americans. Communities and work sites have ongoing programs that target heart health. Exam ples of available programs are presented here.

Federal Government

 Dietary Guidelines for Americans, 2005. The Dietary Guidelines for Americans are published jointly every 5 years by the Department of Health and Human Services and the USDA. The guidelines provide authoritative advice for people 2 years and older about how good dietary habits can promote health and reduce the risk of major chronic diseases. They serve as the basis for federal food and nutrition education programs. Visit

http://www.health.gov/DietaryGuidelines/dga 2005/document/default.htm.

 Dietary Guidelines for Americans, 2010. Provides the most recent dietary guidelines for Americans. Visit http://www. cnpp.usda.gov/dietaryguidelines.htm

 MyPyramid.gov. MyPyramid offers personalized eating plans and interactive tools that help plan and assess healthy food choices based on the Dietary Guidelines for Americans. Visit http://www.mypyramid.gov/.

 Health and Human Services Small Step Adult and Teen program. Information is presented about steps to take to improve health and well-being. Visit http://www.smallstep.gov/.

National Heart, Lung, and Blood Institute (NHLBI)

- National Cholesterol Education Program. The goal of the NCEP is to reduce illness and death from CHD in the United States by reducing the percentage of Americans with high blood cholesterol. Through educational efforts directed at health professionals and the public, the NCEP aims to raise awareness and understanding about high blood cholesterol as a risk factor for CHD and the benefits of lowering cholesterol levels as a means of preventing CHD. Visit http://www.nhlbi.nih.gov/about/ncep/.
- Hearts N' Parks program is a national, community-based program supported by the NHLBI and the National Recreation and Park Association that is designed to encourage heart-healthy lifestyles in communities. Visit http://www.nhlbi.nih .gov/health/prof/heart/obesity/hrt_n_pk/ index.htm.
- Seventh Report of the Joint National Committee on Prevention, Detection, Evaluation, and Treatment of High Blood Pressure. Updated guidelines for hypertension for the health care professional. The report provides new evidence for treating high blood pressure. Visit http://www.nhlbi.nih.gov/guidelines/hypertension/jnc7full . htm

Centers for Disease Control and Prevention

- · Healthy weight program is designed to help individuals achieve and maintain a healthy weight. Visit http://www.cdc.gov/healthyweight/index.html.
- Nutrition program helps individuals follow a healthy diet. Visit http://www.cdc.gov/nutrition/index.html .
- Physical activity program helps individuals learn about recommendations for active living. Visit http://www.cdc. gov/physicalactivity/index.html.
- Cholesterol website provides information about blood cholesterol levels and risk of heart disease. Visit http://www.cdc.gov/cholesterol/.
- Food and Drug Administration, food labeling and nutrition programs. Information is presented about labeling requirements for foods under the Federal Food Drug and Cosmetic Act and its amendments. Included in this is information about the nutrition facts panel, health claims/qualified health claims, and nutrient content claims. Visit https://www.fda.gov/Food/LabelingNutrition/default.htm.

Professional Organizations

- · American Heart Association
 - Heart-Healthy Grocery Shopping Made Simple program provides a tool for consumers to use that simplifies the selection of heart-healthy foods at the supermarket. Visit http://www.americanheart.org/presenter.jhtml?
 identifier=2115.
 - Delicious Decisions program provides heart-healthy recipes online. Visit http://www.americanheart.org/deliciousdecisions/jsp/home/home.jsp ?_requestid= 1884425.
 - Start! program is a new daily walking program that is designed to improve fitness. Visit http://www.startwalkingnow.org/.
 - My Life Check assists individuals in assessing their current health status, and in establishing a customized action plan to improve heart health and quality of life. Visit http://mylifecheck.heart.org/AboutUs aspx?NavID=2&CultureCode=en-US.
- American Dietetic Association website provides food and nutrition information for individuals of all ages. In addition, there is a link for consumers to find a registered dietitian. Visit http://www.eatright.org/.

Other Programs

Many local, regional, and state programs provide information about a healthy lifestyle. One example is the YMCA Activate America program that is designed to create and sustain healthier communities. Ways to locate these programs include asking health professionals in the community, searching the Internet and print media, and calling the local or state public health office.

CONCLUSION

Clinical intervention trials have demonstrated the efficacy of diet intervention, notably by TLC and DASH, for the treatment of elevated LDL-C and blood pressure. These dietary patterns are reduced in SFA and *trans* -fatty acids, cholesterol, and total fat and emphasize increased consumption of fruits and vegetables, whole grains, heart-healthy protein, and reduced-fat dairy. Additional LDL-C lowering is expected with the addition of plant sterols and stanols, unsaturated fatty acids, soy protein, and fiber. Such dietary approaches also improve the metabolic syndrome profile and beneficially affect non-traditional risk factors, such as markers of thrombosis and inflammation, LDL-C particle size, and apolipoproteins. The Mediterranean diet is another treatment strategy to improve CVD risk status.

Many of the dietary factors emphasized by the TLC, DASH, and Mediterranean diets are included in the *Dietary Guide lines* for Americans, 2005, 2010. These guidelines ensure that energy and nutrient requirements are met through foods, although the inclusion of fortified foods and supplements may afford greater dietary flexibility and help some individuals also meet specific nutrient needs or target certain risk factors. However, it is important to carefully evaluate the efficacy of supplements promoted for CVD risk reduction. Certain dietary supplements, such as fish oil and niacin, have shown promising cardiovascular effects, but there is insufficient evidence for others (such as B vitamins and antioxidants). Clinicians should therefore encourage a food-based approach to CVD risk reduction.

Maintaining a healthy body weight is a key component of cardiovascular health. Many studies have demonstrated that dietary modification (specifically calorie restriction) is a key facilitator of weight loss, and modest changes in body weight

(-10 %) are associated with improvements in LDL-C and blood pressure. Of recent interest has been the relative contribution of specific macronutrients to weight loss; however, data from large-scale clinical trials demonstrate that reduction in total calories is the single most important step for weight loss, although macronutrients may induce differential effects on cardiovascular risk factors. Whereas calorie restriction alone is effective for short-term weight loss, interventions that combine diet therapy and physical activity with continued professional support (including behavioral intervention, diet and physical activity counseling) are most effective in the long term. ³²

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CHAPTER 17

Integrative Medicine in the Prevention of Cardiovascular Disease

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KEY POINTS

- Integrative medicine, which blends standard Western and nontraditional medical practices, is used by 38% of adults and 12% of children in the United States.
- Placebo responses, inherent in all conventional and integrative medical therapies, account for as much as
 20% to 50% of clinical outcomes.
- Traditional Chinese medicine, incorporating acupuncture and herbals, reduces elevated systolic blood pressure and oxygen demand to improve supply-demand imbalances and myocardial ischemia. Tai Chi and Qigong, two forms of Chinese energy medicine, frequently involve a component of exercise that also may reduce blood pressure.
- Ayurvedic medicine, originating from India, includes lifestyle changes, particularly dietary and herbal prescriptions, as well as stress reduction through yoga and meditation.
- Chinese and Ayurvedic medicines, with a long history of traditional use, require further study to assess their actions on cardiovascular risk factors.
- Observational studies on singlenutrient dietary supplements suggest reduced cardiovascular risk that has not yet been validated by randomized trials.
- Blood pressure, body mass index, and cholesterol level are improved by a program of yoga incorporating vegetarian diet and stress management with meditation.
- Behavioral and cognitive training, guided imagery, meditation, biofeedback, and

- progressive muscle relaxation appears to be capable of reducing blood pressure.
- Naturopathic medicine incorporates a number of natural practices, such as dietary manipulation and exercise, to prevent disease and to promote healing; other naturopathic practices,

like homeopathy, requiring validation before they can be recommended.

 Chelation therapy, which seeks to lower calcium and to alter atherosclerosis progression, has the potential for serious side effects and is not recommended on the basis of currently available evidence.

Complementary and alternative medicine (CAM) consists of a diverse group of practices and health care systems that generally are not considered to be part of usual Western or allopathic medical practices (Box 17-1). Most academic medical centers in the United States have adopted the term integrative medicine, recognizing the usefulness of combining conventional and CAM practice. There is a large diversity of disciplines that make up CAM therapies (see Box 17-1), and because scientific and clinical studies supporting their mechanisms of action and clinical utility vary tremendously, most academically based centers tend to focus on only a few that are supported by evidence.

This chapter discusses areas of integrative medicine that are supported by evidence (Box 17-2) and that can be used in preventive approaches to cardiovascular particularly coronary artery disease. Some therapies have little rationale or support for efficacy in preventive cardiovascular medicine and are discussed only briefly (Box 17-3). Most guidelines (48%) that have been developed are based on recommendations that are less scientifically rigorous and expert opinion, case studies, or standards of care. 1 Despite the absence high-quality prospective randomized clinical trials on cardiovascular integrative medicine, prevention in experimental studies have identified the mechanisms by which these therapies may reduce cardiovascular risk and, by extension, establish the potential for their clinical action.

In addition to its action on traditional cardiovascular risk factors, the influence of CAM on stress is discussed throughout this chapter because it is linked to many aspects of coronary disease, 2 including blood pressure dysregulation, diabetes, and cholesterol. 3,4 This chapter, therefore, provides a brief overview of available clinical and basic science evidence for the use of integrative approaches cardiovascular disease risk, focusing on our current Western understanding of CAM rather than on traditional CAM theory, which can be found in recent texts. 5-8 Dietary and nutritional approaches are considered in Chapter 16, although a discussion of supplements, including herbals and vitamins as they relate to cardiovascular risk reduction, is provided here.

USE OF INTEGRATIVES MEDICINE IN THE UNITED STATES

The National Center for Complementary and Alternative Medicine (NCCAM) recently released an update on the use of CAM practices in the United States, taken from data collected in the National Health Interview Survey of 2007. 9 Approximately 38% of adults and 12% of children use some form of CAM. CAM is used with higher frequency by women and children of families that use CAM. The most common uses are for pain, anxiety or stress, hypercholesterolemia, and insomnia (Fig. 17-1). A major issue is the absence of communication between patients integrative medical therapies and their physicians, in part because access is obtained outside the standard medical environment and because of the lack of approval of many of these unconventional therapies by the medical



- Whole medical systems (traditional Chinese, Ayurvedic medicine , naturopathy, homeopathy)
- Mind-body medicine (meditation, yoga)
- Biologically based practices (dietary supplements, herbs, foods, vitamins)
- Manipulative and body-based practices (chiropractic, osteopathic manipulation massage)
- Energy medicine (Qigong, Tai Chi, and bioelectromagnetic therapies)

profession. Thus, interest in CAM is driven by the lay community **more** than by the medical community.

Despite the absence of acceptance in the past, there is increasing use and referral by some practitioners, mainly family physicians and other primary care providers, as they are encouraged by their patients, as studies begin to appear, and as many schools begin to introduce CAM education into their curriculum. ¹⁰ It is apparent, however, that there are only a limited number of cardiologists who use or refer patients for CAM therapy and fewer still who study it. 6-8,11 The skepticism of many physicians stems from a common belief that much if not all of the clinical effect of CAM is equivalent to that of a placebo.



- Acupuncture
- Tai Chi, Qigong Chinese herbals
- Ayurvedic medicine
- Yoga
- Meditation
- Ayurvedic herbals
- Naturopathic medicine
- Western supplements

ALTERNATIVE MEDICINE

ROLE OF PLACEBO IN COMPLEMENTARY AND

Placebo, translated from the Latin phrase "I shall please," is part of every clinical intervention, whether standard Western therapy or CAM procedure, and clearly has been well recognized in many forms of cardiovascular medicine. 12-14 Simple interaction between a provider and a patient frequently leads to clinical improvement unrelated to the intervention in many diseases. 15 In fact, between 20% and 50% of response to any medical treatment, including standard Western thera pies for cardiovascular disease, may be ascribed to a placebo effect, either because of a physiological placebo-related response or because of regression to the mean with repetitive testing. 16,17 Furthermore, placebo responses are more likely to occur in trials comparing continuous variables like pain and blood pressure.

Chronic stable angina improves by 30% to 50% 18 and blood pressure can be reduced by as much as 20 to 40 mm Hg by reassurance or placebo interventions. 19 In integrative medicine, the concern is that most if not all of the effect is a nonspecific response to placebo, that is, there is no active intervention. This

belief is reinforced by the fact that many

BOX 17-3 Unproven Integrative Therapies for Cardiovascular Disease



- Chelation therapy
- Homeopathy
- Bioidentical hormones

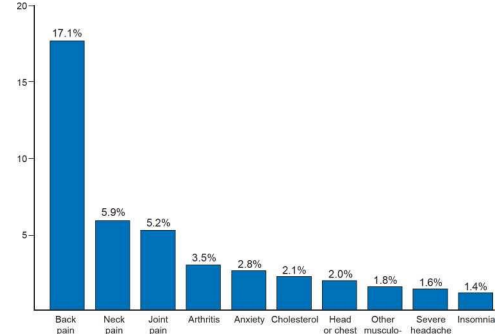


FIGURE 17-1 Diseases and conditions for which CAM is most frequently used among adults, 2007. (From Barnes P, Bloom B, Nahin R: Complementary and alternative medicine use among adults and children: United States,

2007. Natl Health Stat Report 12:1,

2008.)

Diseases/conditions for which CAM is most frequently used among adults-2007

skeletal

274 of the symptoms and diseases treated in cardiovascular medicine, such as angina and stress responses, have improved with placebo therapy. ²⁰ Compounding the difficulty in distinguishing between active and placebo responses is the finding that both operate through the endogenous opioid system, including shared regions of the brain like the periaqueductal gray that are activated by acupuncture and placebo. ²¹⁻²⁵ Fur thermore, unwarranted or exaggerated expectation in patients using CAM interventions may heighten the placebo response. ²⁶ Double-blind controls, placebo interventions, and non-treatment arms have been used to detect placebo responses. 27 It is frequently difficult to truly blind the practitioner, however; for example, the acupuncturist has to know where to place the needle to achieve an optimal response. Furthermore, the use of no or inadequate controls has limited the interpretation of many clinical CAM trials. More than 50% of trials coming from Asian countries are limited by the absence of adequate controls, and once many trials use a suitable control, some studies have found little difference between the sham and the true (verum) intervention, 28

Despite the inherent problems with control intervention in studies of integrative medicine, they are extremely important, and many suitable control interventions have been ¹⁷ devised. One example of a suitable control is in the study of electroacupuncture regulation of elevated blood pressure, in which a needle is placed in an acupoint known to exert biological effects but is not stimulated. Alternatively, needles can be placed in acupoints known not to exert responses. ²⁹ In the first control, afferent nerves are not stimulated, and hence any response is related to interaction between the therapist and the subject. The latter control is associated with sensory stimulation, but neural input to the central nervous system does not involve the areas known to regulate blood pressure. ^{28,30}

Uninformed study subjects cannot differentiate between control and active interventions. ³¹ By use of this paradigm, 30 minutes of low-frequency, low-intensity acupuncture at active acupoints, but not sham acupuncture with either type of control intervention described earlier, lowers elevated blood pressure in experimental studies. ²⁹

TRADITIONAL CHINESE MEDICINE

Traditional Chinese medicine, originating more than 2000 years ago, incorporates several diverse treatment options, including acupuncture and acupressure, Chinese herbals, moxibustion (local heat with a Chinese herb), massage, Tai Chi, Qigong, and dietary therapy. Acupuncture, Tai Chi, and Qigong are forms of energy-based medicine, the energy referred to as *Qi*.

Acupuncture, Acupressure, and Moxibustion

Acupuncture and its derivatives, acupressure and moxibustion, are based on a system of 12 main channels or meridians that lie along the body surface. Along these meridians are small nodes or acupuncture points (acupoints) that direct the therapist where to place the needle to exert pressure or heat. Although neither the meridians nor the acupoints have a physical basis, they are useful because they direct the thera pist to where stimulation should occur.

Mechanism of Action

Many if not all meridians lie over major neural pathways that contain both motor and, more importantly, sensory nerves. Thus, from a physiological perspective, acupuncture needles penetrate the skin and in most circumstances are positioned through underlying nerves or sufficiently near the mixed nerve bundles that contain both sensory and motor fibers. Although stimulation of the muscle motor fibers is not important in the acupuncture effect because paralytic agents do not alter acupuncture's action on the cardiovascular

system, fine muscle contractions are helpful in alerting the practitioner that the needle is positioned in a proper location near the nerve bundle. Conversely, activation of sensory neural pathways provides input to regions of the central nervous system that regulate cardiovascular function. ²⁸

Transection of the afferent pathway central to needle insertion eliminates all but the placebo response to acupuncture . More specifically, acupuncture-related activation of thin-fiber somatic sensory pathways provides strong input to the spinal cord, ventral hypothalamus, midbrain periaque ductal gray, and both pressor and depressor regions that regulate sympathetic (and probably parasympathetic) outflow in the medulla, located in the lower brainstem.

Manual acupuncture or low-frequency, low-intensity electroacupuncture is capable of causing the release of a number of modulatory (inhibitory) neuropeptides in the brain, including opioids (endorphins and enkephalins), y-aminobutyric acid, nociceptin, serotonin, and endocannabinoids as well as excitatory amino acids like glutamate and acetylcholine, that ultimately inhibit sympathetic (and probably parasympathetic) outflow to the heart and vascular system. ^{32,33} In the spinal cord, acupuncture appears to inhibit sensory inflow and sympathetic outflow through both opioid and nociceptin mechanisms of blockade. ^{34,35}

Two important concepts in acupuncture are acupoint specificity and the nature of its action. Point specificity is the differential clinical response to stimulation of specific acupoints. ³²For example, some acupoints, like those along the pericardial meridian overlying the median nerve in the wrist, exert a stronger influence on the cardiovascular system than other points do. ³⁶ The extent of influence is determined by the amount of input to regions of the brain that control cardiovascular function. The nature of acupuncture's action is determined by the mode of sensory nerve stimulation, the duration of stimulation, and the extent of release of neu rotransmitters in the central nervous system.

Low-frequency electrical (2 to 6 Hz) or manual acupuncture sessions for 30 to 45 minutes seems to be most effective, reducing sympathetic outflow after 10 to 15 minutes of stimulation and lasting for many minutes to hours or even days after acupuncture sessions , depending on the model of investigation and extent of repetitive stimulation. Thus, the cardiovascular influence of electroacupuncture can last for 1 to 2 hours in anesthetized experimental animal studies and for 10 to 12 hours in awake animals, whereas repetitive acupuncture in patients can exert an influence on blood pressure for several weeks. ²⁸

Acupuncture's Action on Cardiovascular Risk Factors

A number of cardiovascular risk factors, including hypertension , obesity, and hypercholesterolemia, potentially may be influenced by acupuncture. In addition, there is some evidence that acupuncture may be effective in stroke and coronary as well as peripheral arterial disease. $^{\rm 28}$

Because acupuncture can decrease sympathetic outflow and sympathoexcitatory reflex responses associated with elevated blood pressure, there is a rationale to use it for the treatment of mild to moderate hypertension. ³² However, the results of clinical trials are mixed. Experimental studies in quadriplegic rats suggest that transcutaneous electrical stimulation (TENS), which shares some features of stimulation and physiological response with electroacupuncture, decreases the exaggerated blood pressure responses associated with colon distention. ³⁷ Although acupuncture appears to be safe, ³⁸ no clinical trials are available on its effect in spinal patients experiencing large fluctuations in blood pressure associated

with autonomic dysreflexia. Blood pressure in spontaneously hypertensive rats is reduced by acupuncture at an acupoint located mental stress. 72 The influence of acupuncture is not universal over the deep peroneal nerve for periods lasting up to 12 hours. ³⁹ A because it occurs in only 70% of individuals. ³¹ This raises the small study of 50 patients suggested that 30 minutes of acupuncture question of which individuals are most likely to respond. Whereas lowered both systolic and diastolic pressure. 40 Conversely, the there is no definitive answer to this question, those individuals SHARP (Stop Hypertension with Acupuncture Research Program) demonstrating changes in pain threshold and skin finger trial, which treated patients with moderate hypertension during a temperature in response to acupuncture appear most likely to 12-week period, demonstrated no influence on blood pressure over respond. 73,74 Acupuncture's action on skin temperature signals its and above the response to an invasive sham control when blood action on the sympathetic nervous system, more specifically pressure was measured intermittently with manual mercury cutaneous vasomotor fibers. sphygmo-manometers. 41 However, large and small trials incorporating ambulatory monitoring have demonstrated more decreases nitroglycerin consumption and the rate of anginal attacks consistent decreases in blood pressure in patients with mild to in patients with stable angina. 71,73-75 Finally, a prospective moderate hypertension, especially if acupoints that have been shown nonrandomized study of patients in whom acupuncture I was to have a strong cardiovascular influence (P5, P6, St36, St37, referring administered as part of a lifestyle program incorporating I stress to points along the pericardial and stomach meridians overlying the reduction and healthy eating and living found reduced in-patient median and deep peroneal nerves) are used. 42,43 Acupuncture days, medication use, and accumulated mortality rate. 70,71,76 The appears to influence systolic and mean blood pressure more than independent contribution of acupuncture to 17 these beneficial effects diastolic blood pressure. The onset of action is slow, frequently was not determined. requiring several acupuncture treatments before a sustained decrease in blood pressure is observed. Blood pressure decreases by as well as in patients undergoing reconstructive surgery. 77,78 Spinal 5 to 20 mm Hg and tends to remain low for several weeks after cord stimulation, which may involve stimulation of many of the cessation of treatment.

that acupuncture can lower cholesterol. Daily acupuncture sessions patients with peripheral vascular insufficiency. 80-82 No trials of for a 2-week period reduces the increase in cholesterol in acupuncture's influence in patients with peripheral vascular experimental animal models fed high-cholesterol diets. 44,45 There are disease have been published. no good randomized controlled clinical trials, but a small nonrandomized, unblinded trial of electroacupuncture that did not Tai Chi and Qigong incorporate a control acupoint group demonstrated similar or greater weight loss, low-density lipo protein cholesterol (LDL-C) and triglyceride reductions compared with a control group fed a low-radiate energy, although there is no sound scientific evidence calorie diet. 46

patients provides input to regions of the brain that regulate food ingestion, 47 its ability to assist with weight loss in obesity is less movements of the body, diaphragmatic breathing exercises, certain. Experimental studies ^{48,49} in rats show that auricular (ear) acupuncture leads to a 5% loss in weight during a period of 2 to 3 nervous system, catecholamines, and blood pressure. Chi or Qi weeks. However, clinical trials are mixed, with uncontrolled studies is considered to be energy that helps maintain homeostasis. showing small decreases 49-52 and controlled trials 53-56 showing either According to traditional Chinese medicine theory, this energy very modest or no weight loss that could be attributed to flows through channels called meridians. Although somewhat acupuncture. Many of the trials lacked suitable controls.

Because acupuncture leads to the release of endogenous opioids, it has been thought that it may be useful in treating addictive habits like smoking. In this regard, acupuncture reduces symptoms in input to the central nervous system when they are stimulated. subjects addicted to opiates like morphine . 57,58 However, metaanalyses of relevant clinical trials reveal that many are of low quality, are frequently short term, lack suitable controls, and do not provide taking into account our understanding that the brain (and sufficient information to assess their quality. 59 Thus, at present, spinal cord) strongly controls autonomic and hence insufficient data are available to determine the efficacy of cardiovascular function. 83,84 acupuncture in smoking cessation.

Cardiovascular Responses to Transcutaneous Electrical Stimulation and

Through an opioid mechanism, acupuncture lowers myocardial oxygen demand and hence can reduce demand-supply imbalances and ventricular dysfunction in experimental myocardial ischemia. muscles, to regulate their breathing, and to concentrate their 60,61 Similarly, both TENS and acupuncture reduce myocardial mind. Exercise consists of small postural movements of the ischemia occurring during exercise in patients with angina and limbs, walking, and larger movements. electrocardiographic evidence of ischemia. 62-71 TENS shares some similarities with but is not exactly equivalent to acupuncture because on blood pressure in patients with mild to moderate much higher stimulation intensities and frequencies are used during hypertension. 85,86 For example, two small studies show that the non-invasive TENS stimulation that is not directed at specific locations (acupoints) over neural pathways. Although there is some debate about whether acupuncture can increase **coronary** blood flow, the preponderance of evidence suggests that it mainly reduces ischemia by reducing the increase in blood pressure and double product (but not the elevated heart rate) associated with exercise, hence lowering myocardial oxygen demand. 31

Acupuncture also lowers the reflex excitatory responses to

Application of acupuncture over a course of several weeks

TENS increases the survival of skin flaps in experimental models same central neural systems as in acupuncture, 79 increases skin In addition to hypertension, experimental studies demonstrate temperature and reduces pain, ulcer formation, and tissue salvage in

Energy medicine stems from the belief that all living organisms that demonstrates the existence of bioenergy fields. ² Tai Chi Although stimulation of auricular acupoints to treat overweight and Qigong, belonging to energy medicine, are part of traditional Chinese medicine. Like yoga, they include slow and mental concentration that have an impact on the autonomic controversial, most modern scientists recognize that these meridians, rather than forming a physical entity, represent a road map overlying neural pathways that provide sensory ³² A unique traditional Chinese medicine view is that the physical and emotional hearts are included as a single concept,

> Qigong means Qi training or Qi practice. Of the different forms of Qigong, medical Qigong is most applicable to treat cardiovascular disease and risk factors that promote cardiovascular disease. A typical session of Qigong includes meditation (see later), deep breathing and relaxation, guided imagery, mindful focus, and exercise and is practiced in a quiet place in the fresh air. Thus, practitioners seek to relax their

> A number of studies have evaluated the influence of Oigong

276 Oigong lowers serum catecholamines and blood pressure in patients with essential hypertension. 87,88 However, in general, the quality of these studies is low, and they need to be repeated with larger numbers of patients, better blinding, randomization, and concealment of allocation.

Tai Chi is a type of Qigong that involves meditation, breathing, and slow movements of the limbs. It has roots consumption and weight gain and smaller increments in body processes. 101 mass index and body fat as mea sured by skinfold thickness, groups were small. 91

moderate exercise in patients with mild hypertension. 92 17 hyperlipidemia (preparations of red yeast rice). Decreases in anxiety and in total cholesterol and LDL-C and Danshen elevations in HDL-C, in addition to the blood pressure reduction, have been noted in hypertensive patients participating in a 12week program of Tai Chi. 93,94 However, a review of approximately 70 articles on Qigong for hypertension suggests that most reports are in low-level scientific journals, conference proceedings, book excerpts, and informal reports that were not peer reviewed. Only five studies reported a randomized design. 95

A few studies have investigated the role of Tai Chi and blood pressure similar to that of a music exercise group and showed a greater reduction in diastolic blood pressure. 9

Reiki and therapeutic touch are two other forms of energy medicine. Reiki uses what is believed to be "healing energy" to improve health by inducing deep relaxation. However, there are no studies demonstrating that a practitioner can "transfer showing that there is any direction of energy in therapeutic touch. The responses most likely are simply a "placebo" occurs between the therapist and the patient. 99

Chinese Herbs

Traditional Chinese medicine employs herbs as frequently as acupuncture and frequently in conjunction with acupuncture. Herbs in traditional Chinese medicines are used to correct energy flow and balance as noted earlier for acupuncture and other traditional Chinese medicine therapies. Most commonly, herbs are taken as a fixed formula or in a fixed combination of several herbs. Each individual herb is thought to address a particular imbalance. In the less formal practice of Chinese folk medicine, herbs are used more simply, some what in the manner of Western herbal medicine.

Herbs most commonly used include astragalus (Astragalus mongholicus); danshen, commonly referred to as Chinese salvia (Salvia miltiorrhiza); dong quai (Angelica sinensis); ginger (Zingiber officinale); kudzu (Pueraria lobata); licorice (Glycyrrhiza glabra); lycium (Lycium chinense); Asian ginseng (Panax ginseng); and schizandra (Schisandra sphenanthera) (see Appendix). Herbal preparations may be administered in many forms, including as a tea, tablet, capsule, or decoction; as such, their standardization and use in clinical trials have been challenging. In China today, traditional Chinese medicine is used alongside conventional pharmaceutical treatment in a holistic approach to treatment.

In a recent review 100 of 167 randomized controlled trials on the

in dance as well as in martial arts. Although there are many use of traditional Chinese medicine involving more than 18,000 styles of Tai Chi, it can be modified for physical disabilities. participants, the authors noted that the overall methodological Typically, the degree of exercise stress is considered mild to quality of the trials was rated poor because of lack of description or moderate, increasing heart rate reserve in one study by 58% incorporation of adequate sample sizes, randomization, allocation and oxygen consumption to 55% of peak capacity. 89 Tai Chi and blinding procedures, and disclosure of sample size estimates. increases aerobic capacity to the greatest extent in Description of the quality control of herbal products also has been deconditioned subjects. 90 A small study has shown that lacking from ran domized controlled clinical trials reported in the compared with sedentary age-matched controls, during a 5- literature addressing such issues as quality of the herbs selected, year period, elderly male and female subjects who regularly growing conditions, methods of preparation, testing for heavy use Tai Chi experience smaller decrements in peak oxygen metals and microbial contaminants, and use of good manufacturing

Despite some severe limitations with traditional Chinese although differences between the control and intervention medicine herbal interventions used in clinical trials, a number of compounds have been reported to be efficacious for the treatment of During a 12-week period, Tai Chi mildly lowers systolic cardiovascular conditions (Table 17-1), such as angina pectoris and diastolic arterial blood pressure (-7/2 mm Hg) much like (danshen, compound salvia, suxiao jiuxin wan, tongxinluo) and

According to traditional Chinese medicine, chronic stable angina belongs within the scope of pectoral pain and stuffiness (obstruction of Qi and blood in the chest). The traditional Chinese herbal medicine danshen, obtained from the dried root of Salvia miltiorrhiza , promotes blood circulation and relieves blood stasis. It is one of the most versatile traditional Chinese medicine herbals that has been used for hundreds of years in the treatment of numerous ailments. Much of the early research on the pharmacological actions of Qigong in patients with coronary artery disease. Patients ran danshen has been documented through intravenous use in animal domized to Tai Chi and Qigong 3 weeks after an acute models and human subjects. 102 Because of its properties related to myocardial infarction demonstrated a reduction in systolic improving microcirculation, enhancing coronary vasodilation, and protecting against myocardial ischemia through its negative chronotropic effects as well as suppressing the formation of thromboxane and inhibiting platelet adhesion and aggregation, danshen is widely used either alone or in combination with other herbals.

Danshen has also been studied with respect to its actions on lipid energy" through strategic points (chakras) to cause a lowering (inhibition of LDL oxidation) and hypertension (inhibition demonstrable cardiovascular response. 97,98 In contrast, ther - of angiotensin-converting enzyme). 102 It is indicated for use for apeutic touch, in which the hands are used to direct healing patients with coronary artery disease and other cardiovascular energy, in several studies has demonstrated reduction of diseases in China and to a lesser extent in Japan, the United States, anxiety in coronary care unit patients. 97 There is no evidence and other European countries. In China, danshen is used to treat angina pectoris, hyperlipidemia, and acute ischemic stroke.

The primary active ingredients containing tanshinones and interaction involving reassurance and stress reduction that phenolic compounds in danshen are found in dried root and rhizome preparations. Although danshen has no major side effects, it has the potential to interact with anticoagulants 103 and antiplatelet drugs or supplements with those properties. 104 Hence, it may increase the international normalized ratio (INR) and the risk of bleeding and should be avoided in patients taking warfarin. Danshen also may interfere with serum digoxin measurements. 105 Side effects include pruritus, upset stomach, and reduced appetite. 102

> Evidence to support the use of danshen preparations is too weak for any judgment to be made about its effects. Collectively, evidence from randomized controlled trials is insufficient and of low quality. The first documented systematic review on the quality of randomized trials of danshen was recently published according to the CONSORT standards (Consolidated Standards for Reporting of Trials for

TABLE 17—1 Composition of Popular Traditional Chinese Medicines for Cardiovascular Indications						
Common Name	Latin Name	Active Constituents	Uses	Purported Actions		
Single-ingredient Preparations						
Danshen	Salvia miltiorrhizae	Tanshinones, phenolic	Angina pectoris	Coronary vasodilation		
Red rice yeast	Monascus purpureus	compounds Monocolin K	Lipid lowering	HMG-CoA reductase inhibitor		

Red lice yeast		Monascus purpureus	WOTOCOIIT K	Lipid lowering	THING-COA reductase inhibitor
Compound-ingredient Compound salvia preparations	Prepar	rations		Angina pectoris	
Danshen		Salviae miltiorrhizae	Tanshinones, phenolic		
		Panax notoginseng	compounds		Increased plasma levels of radix chuanxiong
Sanqui		Cinnamomum camphora	Ginsenosides		Increased cardiac contractility
Borneol		Ligusticum chuanxiong Hort			Decrease blood pressure
Suxiao jiuxin wan Radix chuanxiong Borneol		Borneolum syntheticum		Angina pectoris	Anticoagulant
(synthetic					Increase cardiac output
formulation)		Ginseng root, 10%-20%			Increased cardiac contractility,
Tongxinluo		Scorpio, 10%-20%		Angina pectoris	lower blood pressure
Ren shen		Leech, 20%-30%	Ginsenosides		Decrease heart rate
Scorpion		Eupolyphaga tallow steleophage, 10%-			Anticoagulant, coronary vasodilation Analgesic
Leech Ground beetle	20%	Scolopendra, 6%-15%			and sedative
Centinede		30010periura, 0 /0-13 /0			

Slough cicada

Periostracum cicadae, 10%-20%

Traditional Chinese Medicine), which used the Jadad quality scale, for

systematic review of 17 randomized controlled trials in patients with unstable angina noted significant improvement with compounded salvia in combination with standard therapy. Salvia graphic parameters and Root of common peony Borneol (synthetic formulation) Radix paeoniae rubra, 5%-15% Borneolum syntheticum, 2%-10%

Modified from Wu T, Harrison RA, Chen X, et al: Tongxinluo (Tong xin luo or Tong-xin-luo) capsule for unstable angina pectoris. Cochrane Database Syst Rev (4):CD004474, 2006.

were identified, with only 6.7% of the randomized controlled trials the studies was low, thus limiting the clinical application of this being identified as high quality (Jadad score > 4). The authors herbal preparation. concluded that the overall quality of these trials has not improved over time, and the evidence base for danshen is still poorly alternative to standard therapy for the control of anginal symptoms developed. More evidence from high-quality trials is needed to if other standard therapies fail. However, the data are not robust, support the clinical use of danshen preparations. 106

Recommendation. Evidence to support the use of danshen anticoagulants. Recommendation. Evidence to support the use of danshen anticoagulants.

preparations is weak. Collectively, information from randomized controlled trials is insufficient and of low quality. Danshen has the potential to potentiate the effects of warfarin , and as such, it is prudent not to combine use of danshen with any anticoagulant or antiplatelet drugs.

Suxiao Jiuxin Wan

Another popular Chinese herbal that has been used for the treatment of angina pectoris is suxiao jiuxin wan. This preparation may cause remission of angina pectoris, improve anginal symptoms, and reduce the use of nitroglycerin. 110 The ingredients

Sage

and is one of the alternative therapies widely used in China when recent Cochrane Review identified 15 randomized controlled trials long-acting nitrates are not an option. The primary herbal ranging from 4 weeks to 2 years in 1776 patients comparing the ingredients in this formulation include danshen (Salviae effects of suxiao jiuxin wan with nitroglycerin, danshen, and heterogeneity in study endpoints, such as anginal symptoms and outcomes that varied electrocardiographic changes. 108 Methodological improvements with more well defined outcomes are needed. A subsequent

Coronary vasodilation

studies published in China from 1998 to 2007. A total of 150 studies reduced anginal symptoms. 109 Again, the methodological quality of

Recommendation. Compounded salvia may be a viable alternative to standard therapy for the control of anginal symptoms if other standard therapies fail. However, the data are not robust, and caution should be used in patients receiving warfarin or other anticoagulants.

symptoms, and reduce the use of nitroglycerin. 110 The ingredients in suxiao jiuxin wan include radix chuanxiong (Ligusticum Compound salvia has been promoted to improve blood circulation chuanxiong) and borneol (Borneolum syntheti cum) (see Table 17-1). A miltiorrhizae), sanqi (Panax notoginseng), and borneol (Cinnamomum isosorbide dinitrate by evaluating electrocardiographic and angina camphora). 107 The main active component in the compound is endpoints. Unfortunately, the treatment regimen varied danshen (see Table 17-1). A recent meta-analysis supported by the tremendously. In 10 studies, suxiao jiuxin wan provided better Chinese Cochrane Center of 27 randomized trials (n = 3722) to - electrocardiographic results, better improvement in anginal evaluate the effectiveness of compounded salvia preparations symptoms, and less nitroglycerin use compared with patients -(danshen pill) compared with isosorbide dinitrate concluded that the randomized to nitroglycerin alone. Two studies reported electrocar salvia preparations demonstrated a significant improvement in diographic improvements with suxiao jiuxin wan compared with angina symptoms along with electrocardio graphic improvements danshen. Clinical symptoms of angina were also improved with and few adverse events. The adverse event rate was significantly less suxiao jiuxin wan. The one study that compared suxiao jiuxin wan than that of nitrates (2.4% versus 29.7%), leading to greater with isosorbide dinitrate noted no improvement in the withdrawal of patients from the nitrate intervention compared with electrocardiographic or anginal symptoms . As noted before, studies the salvia intervention group. However, there was significant tended to be of poor quality and lacked clinically relevant event

278 between trials. Headaches and bradycardia were reported in some studies, but none required medical management. 111

Recommendation. Suxiao jiuxin wan is not recommended for cardiovascular use until further well-controlled trials with clinically relevant endpoints can substantiate benefits and standard therapies.

Tongxinluo

A new drug studied clinically only since 1995 for its efficacy in have demonstrated that tongxinluo possesses pleiotropic effects that are cardioprotective, including the improvement of endothelial function, lipid lowering, antioxidation, vaso dilation, antithrombosis, anti-inflammation, antiapoptosis, similar to those of statin therapies. 116

In a study using a rabbit model of induced atherosclerosis, tongxinluo in a dose-dependent manner increased the thickness of fibrous caps and plaque contents of smooth muscle cells and collagen; reduced serum lipoprotein levels, inflammatory biomarkers, and mRNA expression of matrix metal loproteinases; and decreased the incidence of plaque ¹⁷ breakage ¹¹⁷ Tongxinluo capsules contain both herbal and insect (scorpion, leech, and centipede) ingredients (see Table 17-1).

A systematic review performed by the Cochrane Collaboration identified 18 short-term studies (15 randomized) luo with or without other treatments (danshen, isosorbide mononitrate, or low-molecular-weight heparin) in patients with unstable angina. The caliber of the studies was rated low, and thus no definitive conclusions were drawn about its efficacy with respect to myocardial infarction, angioplasty, or coronary artery bypass grafting. However, the authors noted adverse events, slight gastrointestinal discomfort and ecchymoses were noted in a few cases. The safety of trials. Additional efficacy studies are necessary for a thorough 104 evaluation of this herbal preparation. 118

Recommendation. The data is preliminary and weak; the safety of tongxinluo remains to be determined. Use of this combination product is not recommended.

Red Yeast Rice

Perhaps the most well known traditional Chinese medicine employed. herbal therapy in Western clinical practice is red yeast rice (Monascus purpureus), used to treat hypercholesterolemia. Conclusion Single herbs touted for lipid lowering include tumeric (Curcuma longa), hawthorn fruit (Crataegus monogyna), coptis or goldthread (Coptis deltoidea), soybean (Glycine max), five-leaf gynostemma (Gynostemma pentaphyllum), green tea (Camellia sinensis), Chinese rhubarb (Rheum palmatum), fleece flower tuber (Polygonum multiflorum), cassia seed (Cinnamomum aromaticum), and Panax ginseng (see Appendix). A number of compound recipes for lipid lowering have been used in recent years and have shown therapeutic effects by improving either clinical signs and symptoms or pathological changes. 119

Red yeast rice is derived from a yeast that grows on rice. It AYURVEDIC MEDICINE has been an Asian food staple and traditional remedy for thousands of years. One of the active ingredients in the yeast, Ayurvedic medicine is one of the oldest systems of natural medicine monacolin K or lovastatin, is an inhibitor of HMG-CoA reductase, the rate-limiting enzyme in the pathway of cholesterol synthesis. The concentration of lovastatin varies in red rice yeast but averages nearly 0.4% by weight.

A meta-analysis of 93 randomized controlled trials (n = 9625) using three different commercial preparations of red yeast rice (cholestin, xuezhikang, and zhibituo) demonstrated a mean reduction in total cholesterol of 0.91 mmol/L, LDL-C of 0.73 mmol/L, and triglyceride of 0.41 mmol/L and a mean rise in HDL-C of 0.15 mmol/L compared with placebo. These levels are comparable to those achieved by many of the standard pharmacological lipid-lowering agents, 120 except for the most powerful statins.

Xuezhikang, a commercial red yeast rice product containing 0.8% reduction of anginal symptoms comparable to that of current lovastatin, or approximately equivalent to 10 mg of lovastatin, has shown impressive results in clinical studies. The China Coronary Secondary Prevention Study carried out in 65 hospitals in China randomized 4870 patients with a prior history of myocardial infarction to either xuezhikang 600 mg twice daily or placebo for a reducing episodes of angina pectoris is tongxinluo. Studies mean duration of 4.5 years. The primary endpoints of nonfatal myocardial infarction and fatal coronary events, cardiovascular mortality, and total mortality were significantly reduced in the treatment group. The need for coronary revascularization was similarly reduced. Total cholesterol was reduced by 13% and LDL-C and enhancement of angiogenesis. 112-115 These effects are by 20%, with a noted 4.2% rise in HDL-C. No treatment-related serious adverse events or deaths were reported during the study period. 13

It is possible that other monacolins or lovastatin hydroxyl acid, plant sterols, isoflavones, and isoflavone glycosides present in xuezhikang also could have cardioprotective effects in addition to that attributed to monacolin K. Reported side effects to red rice yeast are limited but include gastrointestinal upset, headaches, and dizziness. Similar cautions typical of HMG-CoA reductase inhibitors regarding potential side effects, including myopathy, hepatitis, and rhabdomyolysis, should be considered with red rice yeast products and will depend on the levels of monacolin K present. 121 As different involving 1413 subjects that evaluated the benefit of tongxin-concentrations of monacolins exist in an array of red yeast rice products, clinicians should examine the specifica tions of individual products before prescribing. Purity of products may also be variable; some may possess a toxic byproduct of yeast fermentation called citrinin. 122 Future use of xuezhikang as well as of other traditional Chinese recipes for lowering cholesterol will depend on the separation, identification, characterization, and development of some benefit in reduction of angina and electrocardiographic carefully formulated preparations. Additional well-controlled trials improvement. Although there were no recorded severe are needed before clinicians can use these products confidently. Although red yeast rice remains available in the United States, it is now fermented by a different process and the active ingredient has tongxinluo remains to be evaluated in appropriately designed been removed, making its ability to lower cholesterol questionable.

> Recommendation. Red yeast rice products lower choles terol and may be recommended to patients who prefer alternative native treatments or who cannot tolerate standard drug therapies. Choice of products should be carefully scrutinized, and standard laboratory monitoring as used for HMG-CoA reductase inhibitors should be

Collectively, the data on the effectiveness of Chinese herbs in the treatment of angina pectoris is not compelling, except possibly for danshen, which must be used with care when it is combined with conventional medications. Herbs used to treat hyperlipidemia (red yeast rice) are supported by limited data for efficacy. Additional well-controlled clinical trials with standardized preparations in appropriately studied population groups are needed to further define the efficacy and safety of these products.

(originating 4000 to 5000 years ago), older even

medical systems originated from it. 123-124 It originated in India, and lowering, however, have yielded mixed results, with significant the Sanskrit word ayurveda means knowledge or science of life. The design flaws compromising the trials of garlic's effectiveness. Shortmain concept in Ayurveda is living in harmony with the universe term studies have shown some benefit in lipid lowering, whereas (environment) across one's life span. The original Vedic text, long-term studies of 6 months or more fail to show sustained benefit forming the foundation for this system, provides treatment of when garlic is used as a single agent. 136 A recent well-designed disease that includes diet, herbals, lifestyle, and disease prevention. randomized controlled trial using highly characterized diet and -Disease occurs when there is an imbalance between the body and supplement interventions comparing the effects of raw garlic, pow the mind. Treatment, based on mind-body constitutions called dered garlic supplement, aged garlic supplement, and placebo in doshas, uses dietary, lifestyle, and herbal prescriptions to evoke 192 moderately hypercholesterolemic adults demonstrated no natural healing. There is a strong correspondence to traditional significant difference in LDL-C between treatment groups. ¹³⁷ A Chinese medicine and other integrative therapies like naturopathy. systematic review of 21 garlic studies to evaluate the reporting Heart disease, called *hydroga*, is related to several different causes, quality, safety, and efficacy of randomized controlled trials for lipid including, among others, emotional turmoil, dietary indiscretion, lowering demonstrated that 53% of the I garlic trials reported and sedentary lifestyle.

Ayurvedic therapies include yoga and transcendental meditation (a variant of yoga) in addition to dietary alterations and from poor methodology, and results have revealed small, mostly herbals. Yoga means "to join" and includes breath control, physical insignificant decreases in blood pressure. A meta-analysis exercise, and meditation. A 3-month residential yoga program, published in 1994 reported promising results in subjects with mild which includes a vegetarian diet and regular yoga practice, can hypertension but found insufficient evidence to recommend garlic improve a number of cardiovascular risk factors, including blood for clinical therapy. 134 A sub sequent systematic review of 27 small, pressure, weight (body mass index), and LDL-C. 125,126 These randomized controlled trials of at least 4 weeks' duration comparing responses are similar to those noted in small early studies of lifestyle garlic with placebo, no garlic, or another active agent reported modification that incorporated dietary modification, exercise, and mixed effects of various garlic preparations on blood pressure. 136 stress management. 127 Patients with angiographic coronary disease, Two meta-analyses published in 2 0 0 8 139,140 concluded that not receiving lipid-lowering therapy, who practice yoga experience compared with placebo, garlic significantly lowered systolic less angina and improved exercise capacity in association with blood pressure in hypertensive individuals but not in atherosclerosis regression or reduced progression of disease. 128 normotensive individuals. However, the sample sizes of these Small controlled and uncontrolled studies suggest that yoga reduces two meta-analyses of hypertensive patients were not large. blood pressure in patients with hypertension. 129-131 Regular yoga Garlic preparations and doses of 600 to 900 mg/day, providing enhances mood and emotional well-being, indicating improved 3.6 to 5.4 mg of allicin, were common in both meta-analyses. The quality of life. 132 Much of the yoga relaxation response is related to level of blood pressure lowering was comparable to reductions reduced neuro endocrine stress as measured by urinary seen with some antihypertensive drugs, suggesting that garlic catecholamines, dopamine, and aldosterone or by heart rate and may be a nonpharmacological alternative for individuals with skin conductance. 133 The long-term cardiovascular influence of borderline or mild hypertension. However, additional studies yoga has not been adequately studied.

Herbs in Ayurvedic Medicine

The medical system of Ayurveda is popular worldwide, not just in India. Recent analysis of the US National Health Interview Survey 2007 Complementary and Alternative Medicine Supplement to 4 years without reports of toxicity. The concomitant use of estimates that 214,000 adults used an Ayurvedic product in the past 12 months, a 28% increase from the 2002 survey on CAM use. 9 Herbs, minerals, and metals are used in Ayurvedic herbal products. Ayurvedic medicines are divided into two major types: herbal only acid (EPA) may increase antithrombotic effects. As such, and rasa shastra. In the practice of rasa shastra, herbs are combined individuals taking warfarin should be monitored more closely if with metals, minerals, and gems and appear to have been used they are consuming garlic supplements. 104 Additive effects on safely for centuries when they are properly prepared. A number of the more popular Westernized Ayurvedic preparations with randomized controlled trials have been evaluated for efficacy, quality, and product effectiveness . Among these preparations are vascular risk reduction for lowering blood pressure. It is not garlic (Allium sativum) for hyperlipidemia and hypertension and recommended for lowering cholesterol. Consumption of dietary guggul (Commiphora mukul) and arjuna (Terminalia arjuna) for sources of garlic is safe and can be incorporated into a hearthyperlipidemia.

Garlic (Allium sativum)

Garlic has long been touted as a natural product useful for modulation of immune system activity, treatment of hyper lipidemia and hypertension, and primary and secondary prevention Association notes that garlic has "no major role" in lipid lowering of myocardial infarction. Allicin, the bioactive component (total cholesterol and LDL-C). 141 responsible for the cardiovascular activity of garlic, is rapidly 280 Guggul (Commiphora mukul) formed from the allyl sulfur compounds (such as alliin) in the garlic. Raw crushed garlic has the

highest concentration of allicin. Multiple mechanisms of 279 action have been proposed, including decreases in cholesterol and fatty acid synthesis and cholesterol absorption as well as potent antioxidant properties.

Epidemiological studies have shown an inverse correlation between garlic consumption and a reduced risk of cardiovascular

than traditional Chinese medicine because Chinese and other disease progression. 134,135 Clinical studies of garlic's efficacy in lipid positive efficacy with a mean safety score of 63 of 100. 138

Studies of garlic's effectiveness in hypertension have also suffered with adequate sample size and standardized garlic preparations are needed to confirm these findings. Evidence for supplementation with garlic for either primary or secondary prevention of heart disease is not sufficient for its use to be recommended for hypertension.

Garlic preparations have been used in clinical studies for up garlic with herbs or drugs that have warfarin constituents or affect platelet aggregation could increase INR and risk of bleeding. Garlic combined with fish oils and eicosapentae noic cholesterol lowering have been seen when garlic is taken with prescription drugs.

Recommendation. Garlic at best may offer modest cardio healthy dietary plan. Pharmacological treatment guidelines should be followed for individuals identified with car diovascular risk factors. Garlic as adjuvant therapy may be an option for some patients, although the American Heart

Guggul is the gummy resin derived from the bark of the mukul myrrh tree in India but can also be found in countries extending from northern Africa to central Asia. In fact, the mukul myrrh tree has been placed on the Red Data List 142 for further evaluation as it is on the threatened list in two regions in India where it is found because of excessive harvesting. It has played a role in Ayurvedic medicine for several thousand years and, in addition to its cardiovascular role, is used in the treatment of arthritis

and digestive, skin, and menstrual problems.

to be due to its multiple pharmacological activities, notably the hypolipidemic, antioxidant, and anti-inflammatory effects. Gugulipid is the ethyl acetate extract of the gum containing 4.09 Z- and E- guggulsterones per 100 g. The lipidlowering effect of gugulipid and guggulsterone, the bioactive constituent of guggul, has been consistently demonstrated in a number of animal species.

Guggulsterone has been identified as an antagonist to the farnesoid X nuclear receptor (FXR), 143,144 a key transcriptional clinical regulator of cholesterol and bile acid homeostasis. 145-147 More antihyperlipidemic, antioxidant, and anticoagulant properties. recently, guggulsterone has also been shown to be an antago- 17 receptors and an agonist of the pregnane X receptor (PXR), progesterone, and estrogen receptors (ER a). 148,149

Clinical studies of guggul have demonstrated significant reductions in total cholesterol and LDL-C of 15% to 23% and triglyceride reduction of 20%, but results have been variable because populations under study have had different ethnic existing hyperlipidemia. The largest clinical study to date with 205 hypercholesterolemic or hypertriglyceridemic patients was conducted in 1989. After an 8-week diet and placebo lead-in, patients were randomized to gugulipid or placebo daily for 12 weeks. Total serum cholesterol was decreased serum cholesterol by 11% and triglycerides by 17% compared with clofibrate reductions of 10% and 22%, levels by 60%. A longer term study (24 weeks) administering 100 mg of gugulipid daily, as guggulsterone, in conjunction safety profiles with dietary modification also has shown significant recommended. reductions in lipid levels in hypercholesterolemic patients but no improvement in HDL-C. 151 In a randomized placebo- Conclusion controlled trial of gugulipid in the United States, 103 healthy adults with hypercholesterolemia were given 1000 mg (low dose) or 2000 mg (high dose) gugulipid containing 2.5% guggulsterones for 8 weeks. Patients treated with gugulipid experienced no improvement in their lipid levels. In fact, their LDL levels increased by 4%, whereas patients who received placebo experienced a 5% decrease in LDL levels. The effects of gugulipids on HDL were mixed. Also noted was a median serum high-sensitivity C-reactive protein level that was decreased by 29% in the high-dose gugulipid group, whereas it increased by 25% in the placebo group, thus suggesting that gugulipid may also possess anti-inflammatory effects. A hypersensitivity rash was reported in a small number of subjects. ¹⁵² A systematic review of seven guggul studies (133 subjects) to evaluate the reporting quality, safety, and efficacy of published randomized controlled trials for lipid lowering concluded that 86% of the guggul trials reported positive efficacy with a mean safety score of 71 of 100. 138 It appears that individuals on a Western-style diet may achieve less of a lipid-lowering effect from gugu- lipid compared with those consuming a more traditional Indian diet.

Currently, no clinical studies have been conducted to evaluate the safety of long-term use of guggul or guggulsterone, but gugulipids have been shown to cause gastrointestinal upset, headache, mild nausea, belching, hiccups, and rash, depending on the dose and formulation (gugulipid versus raw guggul). Currently, 2.5% guggulsterone content is the minimum standard for quality gugulipid preparations. 153 Concomitant oral administration can reduce propranolol and diltiazem bioavailability and hence may reduce the therapeutic effects of these drugs. 154 Guggul also may have antiplatelet effects. 121

Recommendation. Data on the efficacy of guggul for the

treatment of cardiovascular risk factors are mixed and limited. At The cardiovascular therapeutic benefits for guggul appear this time, guggul is not recommended as a substitute for standard pharmacological therapies for lipid lowering.

Terminalia arjuna

This is a deciduous tree found throughout India. Its bark has been used in Ayurvedic medicine for more than three centuries, primarily as a cardiac tonic. Arjuna is purported to be useful in alleviating anginal pain, hypercholesterolemia, heart failure, and coronary artery disease. It has been shown in animal studies and to have cardiotonic, trials antihypertensive

Among the bioactive constituents in arjuna are tannins, nist at the mineralocorticoid, glucocorticoid, and androgen triterpenoid saponins, flavonoids, gallic acid, ellagic acid, oligomeric proanthocyanidins, phytosterols, and several minerals (calcium, magnesium, zinc, and copper). 155 The antioxidant cardioprotective effects are attributed to flavonoids and oligomeric proanthocyanidins; the positive inotropic effects may be caused by saponin glycosides. 156

A systematic review of six studies of Terminalia arjuna backgrounds, dietary habits, body weight, and severity of incorporating 390 patients to evaluate the reporting quality, safety, and efficacy of published randomized controlled trials for lipid lowering demonstrated that 100% of the trials reported benefit and had a mean safety score of 20 of 100. 138 Terminalia arjuna has been shown to be efficacious in a small randomized clinical trial to evaluate the effectiveness of an extract in patients with stable angina lowered by 24% and triglycerides were decreased by 23% in compared with placebo and isosorbide mononitrate. 157 The arjuna 70% to 80% of patients taking gugulipid. 150 In a subsequent preparation used in these studies was a commercial herbal-mineral 12-week crossover study of 125 patients taking gugulipid compound containing more than eight different herbs, making it compared with a comparable dose of clofibrate, gugulipid unclear whether the cardioprotective benefit was due solely to arjuna.

Recommendation. Data for Terminalia arjuna are limited to a respectively. Unlike clofibrate, gugulipid increased HDL single product formulation evaluated in a small number of subjects. Additional well-controlled comparative trials are needed, and safety profiles must be established before its use can be

Several studies have shown poor quality control of some Ayurvedic medications, particularly some rasa shastra Ayurvedic medicines containing high levels of lead, mercury, and arsenic. 158 More than 80 cases of lead poisoning associated with Ayurvedic medicine use have been reported world wide since 19 78. 159,160 A recent random sample of commercially prepared Ayurvedic medicines purchased from the Internet had high metal concentrations determined by xray fluorescence spectroscopy. One fifth of both the US- and Indianmanufactured medicines contained detectable lead, mercury, or arsenic exceeding at least 1 regulatory standard for acceptable daily metal intake. The presence of metals in non- rasa shastra medicines is also of concern because 17% of these have been determined to have higher levels of metals, the consequence of environmental contamination of the herbs or incidental contamination during manufacturing. 158 Rasa shastra experts claim that these medicines have been used

for centuries and, if properly prepared and administered, are safe and therapeutic. 161,162

Collectively, the data on the effectiveness of Ayurvedic herbs in the treatment of hyperlipidemia or hypertension are not compelling. Additional well-controlled clinical trials with standardized preparations in appropriately studied population groups are needed to further define the efficacy and safety of these products.

MEDITATION AND STRESS REDUCTION

During meditation, there is concentration on a word or a phrase, practices that support normal function. called a mantra. Meditation reduces blood pressure, heart rate, rehabilitation programs that include emotional support can reduce naturopaths, has little documented value for cardiovascular disease. 17 anxiety and improve mortality by up to 25%. 170-173 One phase of area are small, frequently inadequately controlled, and often not and almonds to reduce LDL-C, soy-based foods, and the blinded.

Substantial research has examined the effects of lifestyle modification involving stress relaxation on blood pressure. These studies have used a number of different methods, such as autogenic training, cognitive and behavioral therapy, guided imagery, meditation, biofeedback, progressive muscle relaxation, and yoga. There is substantial overlap between the various techniques of relaxation. Autogenic training, for example, includes focusing on a physiological sensation like the heartbeat and self-suggestion, which can also be used during meditation. Cognitive therapy involves control of irrational thought processes; behavioral reinforcement rewards certain behaviors that promote relaxation. Guided imagery focuses on calming images. A review of more than 25 randomized trials suggests that relaxation lowers systolic and diastolic blood pressure a small amount, ranging from 2 to 8 mm Hg. 176 However, many trials have been poorly constructed, and the transient lowering of cholesterol, are unproven. Because use of so many different techniques to lower stress makes comparison difficult. Furthermore, 15 of the trials that compared mobilization of serum calcium is not likely to be effective in altering relaxation with sham demonstrated an insignificant decrease in atherosclerotic progression. blood pressure. More rigorous evaluation of this commonly included aspect of lifestyle modification is warranted.

NATUROPATHIC MEDICINE

Naturopathic medicine is both a health system and a philosophy -(Box 17-4). It originated from a number of traditional medical poorly designed. 185 The majority of the studies of coronary or systems from India (Ayurvedic), China (traditional Chinese peripheral arterial disease are case reports and cases medicine), Greece (Hippocratic), and Germany (homeopathy, hydrotherapy), among others. It is largely based on the belief that the body can heal itself and thus subscribes to the healing power of nature. The goal is to live as naturally as possible and to use healing practices that support normal physiological function rather than drugs that

Consuming natural unrefined and organically grown foods

Enough exercise

Living a lifestyle that is in moderation

Thinking constructive thoughts and emotions

Avoiding environmental toxins

Maintaining proper elimination

would be considered an artificial enhancement. Naturopathic physicians believe that most (not all) disease is the result of violation of natural laws of living. Thus, healing results from a variety of

Naturopathy consists of health maintenance, disease prevention oxygen consumption, and plasma cortisol as subjects relax. 163-166, education, and self-responsibility. The practice of naturopathy Likewise, meditation in elderly subjects with congestive heart relies on diagnosis and the use of natural therapies that evoke the failure is associated with reduced catechol amine production and body's endogenous healing mechanisms. Naturopaths function as improvement in the quality of life. 167 Depression, which primary care physicians; because they realize that not all diseases predisposes to poor outcome after myocardial infarction, due in part can be prevented or treated by the natural approach, practitioners to low adherence to a low-fat diet, regular exercise, and stress will employ office surgery, prescription drugs, and referrals as management, 168 can be reduced by mindfulness-based stress appropriate. Clinical nutrition, botanical medicine, acupuncture, reduction. 123 However, cognitive therapy for these patients may not hydro therapy, physical medicine, and counseling form the most improve survival after a myocardial infarction. 169 Conversely, compelling therapies. Homeopathy, although practiced by most

Clinical nutrition uses the diet as therapy and in the prevention deep relaxation that has been called the fourth state of of disease. It is the foundation of naturopathic practice. Botanicals, consciousness, to distinguish it from waking, dreaming, and including herbals, can be used in place of some drugs, but they are sleeping, is characterized by reduced plasma cortisol and lactate more commonly used to support natural healing of the body. In fact, levels, decreased metabolism, reduced breathing, increased brain nutritional training in naturopathy generally is much greater than alpha wave activity, and increased cerebral blood flow. ^{174,175} Many that received by allopathic students. Other chapters in this textbook publications come from international journals or American CAM provide information that would be recommended by naturopathic journals with relatively low impact factors because studies in this practitioners, including dietary fiber, omega fatty acids, walnuts Mediterranean diet. 177-180

Chelation Therapy

Chelation therapy with disodium ethylenediaminetetraacetic acid (EDTA), which is used by some naturopathic practitioners, has clear clinical value in treating lead intoxication. It has also been used "off clinical value in treating lead intoxication. It has also been used "off label" for more than 30 years to treat atherosclerosis. Chelation therapy with EDTA, supplemented with some vitamins, heparin, and magnesium, has been promoted by the American College for Advancement in Medicine. EDTA binds divalent and trivalent cations, including calcium. Proposed mechanisms of action, including lowering of calcium from plaques, inhibition of platelet aggregation, mobilization of parathyroid hormone to remove calcium, vascular antioxidant effects including chelation of iron, and calcification of plaques tends to occur late in plaque development,

EDTA infusion carries a risk of congestive heart failure because EDTA is administered in large volumes as the diso dium salt. Renal toxicity, hypoglycemia, and hypocalcemia and tetany can also occur if the infusion rate is too rapid. ¹⁸¹ There is also evidence that EDTA may be a pro-oxidant. 182 Studies show that 100,000 to 800,000 individuals in the United States have received this therapy. 183,184

Clinical trials of chelation therapy are few, small in size, and

282 studies, most uncontrolled and retrospective. One large study and a smaller follow-up study of patients with peripheral vascular disease found no difference between control and treatment groups in symptomatic relief 186 or in angiographic disease and transcutaneous oxygen tension. 187 Three prospective randomized trials of chelation therapy have been published . A Danish trial of 153 patients in 1992 reported no long-term therapeutic effect at 3 or 6 months; a small trial of 32 patients from New Zealand showed some improvement in patients with peripheral vascular disease at 3 months. 186-190 Both trials had significant methodological deficiencies that preclude accurate conclusions. A final 6-month prospective study called the PATCH trial (Program to Assess Alternative Treatment Strategies to Achieve Cardiac Health) of 39 patients with well-documented coronary disease observed no significant difference in clinical outcome or quality of life score. 190 Unfortunately, this study was underpowered, and the placebo arm s contained vitamin C, which may have provided an anti- oxidant effect. A larger randomized trial to assess chelation therapy (TACT) is ongoing. It is unclear why NCCAM funded this \$30 million trial without definitive mechanistic evidence for the action of EDTA disease. 194-197 The metabolism of homocysteine requires several B infusion or support from previous clinical trials.

angina. 191

NUTRITIONAL SUPPLEMENTS

use of herbals, commonly ginseng, as tabulated from the 1999-2002 dilation by only 1.08% compared with placebo. 203

warranted for individual use.

care. Supplements in this category include α -lipoic acid, L-arginine, or secondary prevention of cardiovascular disease. ²⁰⁵ Le carnitine, nonprescription niacin products, and selenium (see Appendix). Dietary supplements continue to be popular as recommendation of folic acid and other B vitamin supplements for documented by the spending of more than \$23.7 billion by primary or secondary prevention of cardiovascular disease. ²⁰⁵ health are shown in Table 17-2.

B Vitamins

Moderate elevations of plasma homocysteine levels have been associated with an enhanced risk for atherosclerosis

272 225 125	21 17 9	
125		
	9	
109	8	
104	8	
58	4	
41	3	
39	3	
rotene 31	2	
	58 41 39	58 4 41 3 39 3 protene 31 2

326

Fish/animal oils

vitamins as cofactors, specifically vitamin B 6, vitamin B 12, and Recommendation. Given the lack of firm evidence supporting a folate. Homocysteine levels can be decreased by the administration beneficial effect of chelation therapy and the potential for serious of supplemental folate, with or without vitamin B 6 or B 12. Although side effects, the American College of Cardiology and the American epidemiological studies suggest potential cardiovascular benefit Heart Association have not endorsed its use in chronic stable with B vitamin supplementation, most intervention trials (such as HOPE-2, VISP, and NORVIT) using combined B vitamin therapy of folic acid and vitamins B 6 and B 12 have shown no benefit. 198-200

More recently, results from the Women's Antioxidant and Folic Acid Cardiovascular Study, a secondary prevention trial of 5442 women, revealed that combination B vitamin therapy, including 2.5 It is estimated that 52% of adults participating in the 1999 2000 mg folic acid, 50 mg vitamin B₆, and 1 mg vitamin B₁₂, after 7 years National Health and Nutrition Examination Survey (NHANES) had of treatment and follow-up demonstrated no reduction in total consumed some form of a dietary supplement within a month cardiovascular events compared with placebo. 201 A meta-analysis before participating in the survey; 35% took a multivitamin- of four randomized, controlled trials of B vitamin therapy found no multimineral supplement. 192 Dietary supplement use in individuals evidence that B vitamin supplements slowed the progression of with coronary artery disease or risk factors for coronary artery atherosclerosis. 202 However, it has been postulated that folic acid disease is common; 60% of adults with coronary artery disease, supplementation, through lowering of homocysteine levels, may stroke, hypertension, or ele vated cholesterol used at least one positively influence vascular function in the early stages of dietary supplement within the month preceding the survey, cardiovascular disease by modulating endothelial dysfunction. A Individuals with a history of coronary artery disease or stroke recent meta-analysis of 14 intervention trials in which high doses of commonly report the use of vitamin E, folic acid, and niacin; those folic acid were administered for 4 months or more and flow was with a history of hypertension or elevated cholesterol report higher measured demonstrated that folic acid improved flow-mediated

A protocol for a collaborative meta-analysis of homocystein -A number of popular dietary supplements with a significant - lowering trials for prevention of vascular disease has been proposed evidence base lack public health recommendations for primary or by the B-Vitamin Treatment Trialists' Collabo ration. By pooling the secondary prevention of cardiovascular disease. These include B data from 12 B vitamin supplementation trials for the prevention of vitamins, the antioxidant vitamins C and E, vitamin D, and coronary heart disease and stroke, it is hoped that more definitive magnesium. On occasion, as appropriate, these supplements may be recommendations on the use of folic acid and vitamin B 12 for the prevention of cardio vascular disease will emerge. Currently, seven Other nutrient supplements purported for cardiovascular studies with 33,755 subjects are being monitored for major coronary disease or prevention that have been less well studied in clinical events. 204 Until such time, the American Heart Association notes trials, or for which there is a lack of data from the United States, that the available evidence is inadequate for the recommendation of remain alternative therapies and should be integrated with great folic acid and other B vitamin supplements as a means of primary

Recommendation. Current evidence is inadequate for the Americans on products in 2007. The most popular products for heart However, they can be used to lower plasma homo cysteine levels in high-risk subjects.

Antioxidant Vitamins

Despite a large body of epidemiological evidence suggesting a favorable association between a diet high in antioxidants and reduced risk of coronary heart disease, no clinical trial has confirmed such benefit. Higher overall intake of vitamin C was associated with lower rates of coronary disease in some cohort studies with information on supplemental vitamin C intake ²⁰⁶ but not in others. ²⁰⁷ Improved endothelial function has been observed with vitamin C supplementation in patients at risk for cardiovascular disease. ²⁰⁸ An inverse relationship between vitamin E intake and the risk of coronary disease demonstrated that 30 IU/day of vitamin E potentially can lower coronary disease risk by 4%. Supplementation with vitamin E alone has also shown an inverse correlation with coronary heart disease. 207

Randomized trials have evaluated antioxidant supplements in varying doses for lowering of coronary heart disease risk. In the Heart Protection Study, 20,536 patients with coronary artery disease or diabetes were randomized to antioxidant vitamins (600 mg of vitamin E, 250 mg of vitamin C, and 20 mg of beta carotene) versus placebo. Although the vitamin regimen was found to be safe, there was no evidence for a therapeutic effect after 5 years of treatment. ²⁰⁹ Similarly, the Physicians' Health Study follow-up also found that neither vitamin C nor vitamin E reduced the incidence of major cardiovascular events, myocardial infarction, stroke, or cardio vascular mortality after 8 years of study. Furthermore, use of vitamin E was associated with an increased risk of haemorrhagic stroke. 210

In contrast, the Antioxidant Supplementation in Athero sclerosis Prevention (ASAP) study in hypercholesterolemic Recent literature suggests that vitamin D inadequacy is pan patients randomized to twice-daily supplements of 136 IU of vitamin E, 250 mg of slow-release vitamin C, both, or placebo demonstrated that only combined supplementation of vitamin E and vitamin C slowed the progression of common carotid intimamedia thickness (25% decrease). 211 However, in the Women's 224-227 and metabolic syndrome, 228-230 as well as in subjects with Angiographic Vitamin and Estrogen study, post menopausal women with some degree of coronary stenosis who took vitamin E and vitamin C twice a day or placebo for more than 4 years experienced no cardiovascular benefit, and in fact, all-cause mortality was significantly higher in women taking the supplements. 212 Whereas this latter observation may have been a chance finding, it follows a trend towards an increase in mortality that was observed in the Heart Protection Study. 209

Randomized clinical trials also cast doubt on the efficacy of vitamin E as monotherapy to prevent coronary disease. Although the Cambridge Heart Antioxidant Study (CHAOS) demonstrated a reduction in events of the combined endpoint of death or nonfatal myocardial infarction by 47%, 213 HOPE and HOPE-2 provide strong evidence that moderately high doses of vitamin E supplements do not reduce the risk of serious cardiovascular events among men and women with established heart disease or diabetes. In fact, participants taking vitamin E in the HOPE-2 trial were 13% more likely to experience and 21% more likely to be hospitalized for heart failure, an unexpected and statistically significant finding. 214 Possible explanations for this harmful effect with cardiovascular disease. Unlike the data and association of of vitamin E could relate to the potential for a -tocopherol to vitamin D with osteoporosis, the cardiovascular data are mixed become a pro-oxidant in an oxidative environment, thereby depressing myocardial function, 215 or to the higher doses than those normally consumed in the diet employed in the study.

Data from randomized controlled trials in women have provided much needed information for clinicians to make appropriate therapeutic choices in the management of cardio vascular risk in women. The Women's Heath Study of 39,876 apparently healthy women aged 45 years and older found that vitamin E (600 IU on alternate days) reduced cardio vascular death, a secondary endpoint, by 24% but had no

effect on total cardiovascular events, myocardial infarction, 283 or stroke. ²¹⁶ In further analysis of the same data, women ran domized to vitamin E demonstrated an overall 21% reduced risk of venous thromboembolism. The observed risk reduction was

44% among those with prothrombotic mutations or a personal history of venous thromboembolism. Overall, vitamin E was associated with a lower risk of bleeding than that observed for low-dose aspirin. 217

The American Heart Association concluded in their 2007 update of the women's prevention guidelines that antioxidant vitamin supplements should not be used for the primary or secondary prevention of cardiovascular disease. 218 However, as observed in the Women's Heath Study, there may be certain populations, such as the elderly (> 65 years) or those with a history of venous embolism, that could benefit from targeted therapy.

A number of meta-analyses have evaluated a range of car diovascular disease outcomes for varying forms and doses of vitamin E in a number of populations of patients. 219-223 Most analyzes concluded that vitamin E provided no benefit and I no increased risk. Lacking from most antioxidant intervention trials are plasma measures of vitamins or determination of a subject's baseline oxidative stress status. Future studies should determine whether those with marginal vitamin status 17 based on measurements will benefit from supplementation.

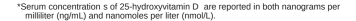
Recommendation. The American Heart Association does not recommend the use of antioxidant vitamin supplements for the primary or secondary prevention of cardiovascular disease. Patients should be encouraged to supplement their diet with foods rich in antioxidants and to consume a multi- vitamin-mineral supplement if they are concerned that their diets are inadequate.

Vitamin D

demic, and a growing body of evidence is emerging to show that low levels of vitamin D may adversely affect the cardio vascular system. In this regard, data are accumulating in populations with elevated cardiovascular disease risk factors, such as hypertension established disease, such as diabetes, 231,232 coronary heart disease, ²³³ peripheral arterial disease, ²³⁴ stroke, ²³⁵ congestive heart failure, ^{236,237} and cardio vascular mortality associated with low vitamin D. ²³⁸ In 1739 participants in the Framingham Offspring Heart Study, researchers found that those with vitamin D blood levels < 15 ng/mL had twice the risk of a cardiovascular event in the next 5 years compared with those with higher levels of vitamin D. 227 Interestingly, the risk remained significant after adjustment for traditional cardiovascular disease risk factors. Calcium and vitamin D supplementation was found neither to increase nor to decrease coronary or cerebrovascular risk in generally healthy postmenopausal women participating in the Women's Health Initiative. 239

Low circulating levels of 25-hydroxyvitamin D are also associated with increased evidence of inflammation, oxidative burden, and cell adhesion, suggesting that vitamin D plays a role in processes that contribute to cardiovascular risk and mortality. ²⁴⁰ Unfortunately, no dose-response studies with supplemental vitamin D are available yet to test the association of vitamin D and less robust and thus lack the evidence base to suggest causation. It is premature to recommend screening for at-risk individuals and supplementation with vitamin D above the recommended levels.

There is considerable discussion of the appropriate serum concentrations of 25-hydroxyvitamin D associated with deficiency versus optimal overall health (Table 17-3). The current



level of variability in 25-hydroxyvitamin D measurements calls into risk for hypertension compared with those in the lowest quintile. question the stability of 25-hydroxyvitamin D assays and their 246 A recent review of nutritional effects on blood pressure indicated ability to accurately reflect the vitamin D status in individuals. ^{241,242} that high levels of magnesium as well as potassium, calcium, and

use of vitamin D for conditions of cardiovascular disease. Current Review evaluating the effect of combined calcium, potassium, and recommendations set by the Institute of Medicine are 400 IU/day magnesium supplementation for management of primary for individuals 50 to 70 years and 600 IU/day for individuals 70 hypertension in adults failed to find robust evidence to support a years and older. Newer clinical trials conducted in healthy adults role for these supplements in the treatment of hypertension. ²⁴⁸ suggest that a dose as high as 10,000 IU/day has been used without ill effects, but the upper level of intake established by the Institute supplementation for the treatment of cardiovascular risk factors. of Medicine of 2000 ÎÛ/day is considered safe. ²⁴³ Although vitamin However, the Joint National Committee on Prevention, Detection, D supplementation above recommended levels given in clinical Evaluation, and Treatment of High Blood Pressure states that diets trials has not shown harm, most trials have not been designed to that provide plenty of magnesium represent positive lifestyle assess harm. 244 In the Women's Health Initiative, increased risk of modifications for individuals with hypertension. 249 General hypercalcemia and hypercalciuria was not clinically significant, but multivitamin-mineral supplements often provide approximately it was associated with a 17% increase in the risk of kidney stones. 25% (100 mg) of the daily recommended need for magnesium, an

for prevention or treatment of cardiovascular disease. However, controlled feeding study in a group of healthy postmenopausal supplementation with 400 to 600 IU/day and no more than 2000 IU women suggested that consumption of this amount of dietary of 1,25-dihydroxyvitamin D₃ for low circulating levels of vitamin magnesium daily for 58 days was inadequate as shown by the onset D or those with poor dietary intakes and low sun exposure can of cardiac arrhythmias. ²⁵⁰Supplementation with the recommended safely be recommended for patients with cardiovascular disease.

Magnesium

Magnesium is cardioprotective. Aging, decreases in dietary intake, and polypharmacy coupled with decreases in intestinal magnesium absorption and increases in urinary magnesium losses place older adults at risk for magnesium deficiency. According to the US Department of Agriculture, less than 60% of adult men and women have an adequate dietary intake of magnesium. 245

Trials of magnesium therapy in acute myocardial infarction have produced mixed results, and chronic use has focused on the role of magnesium as an antihypertensive. A review of more than 28,000 women enrolled in the Women's Health Study indicated that those in the highest quintile of magnesium intake had a decreased

No clinical practice guidelines have been issued regarding the soy seem to have a beneficial effect on hypertension. ²⁴⁷ A Cochrane

There is insufficient evidence to recommend magnesium amount considered safe but that may not be adequate to sustain Recommendation. Vitamin D is not recommended at this time magnesium stores in the body for at-risk individuals. A small welldietary allowance (420 mg for men and 320 mg for women) of magnesium is typically safe in individuals without compromised kidney function but may cause diarrhea, depending on the formulation.

> **Recommendation.** Dietary supplementation of magnesium is not recommended for treatment of cardiovascular risk factors but can be recommended in the form of a general multivitamin-mineral supplement for those with consistently poor dietary intakes.

Conclusion

In general, clinical trials have not provided sufficient evidence supporting the routine use of antioxidants, B vitamins, or other nutrient supplements to prevent cardiovascular disease or to reduce morbidity and mortality. Some investigators have suggested that understanding the potential utility of vitamin E and

vitamin C in the prevention of coronary disease might require longer studies in younger participants taking higher doses of the supplement. ²⁵¹ Further research is needed to determine whether dietary supplements have any protective value for younger, healthier people without risk factors for coronary heart disease.

NON-VITAMIN, NON-MINERAL SUPPLEMENTS

Nearly 18% of participants in the National Health Interview Survey revealed that the most common CAM therapies used were nonvitamin, nonmineral natural products, including fish oil or omega-3 or docosahexaenoic acid (DHA), glucose amines, echinacea, flaxseed oil or pills, and ginseng. ⁹ Among the dietary supplements with possible cardiovascular benefit not endorsed by national public health recommendations but with a developing base of evidence on which randomized clinical trials can be planned or guidelines for use might be formulated include coenzyme Q10, French pine bark, horse chestnut seed extract, and *Ginkgo biloba*. as appropriate,

these supplements can be integrated into clinical practice and are the data with regard to CoQ10 therapy in patients with congestive discussed here. Other supplements purported for cardio vascular heart failure have reached mixed conclusions about efficacy. Of disease prevention that have been less well studied in controlled more interest are the reduced plasma or serum levels of CoQ10 clinical trials or that lack data in US population groups remain that have been documented in observational studies and alternative therapies and should be integrated into medical practice randomized clinical trials in patients receiving statin therapies. ²⁶²with great care. Supplements in this category include artichoke leaf, ²⁶⁶ Decreases in blood CoQ10 levels with statin treatment may be fenugreek, green tea, gugulipid (described earlier), and policosanol related to reduced synthesis as well as to a decrease in circulating for lipid lowering, and hawthorn for angina pectoris and mild levels of CoQ10, as it is carried in the blood by LDL-C. 267 A recent heart failure (see Table 17-1).

Because nonvitamin, nonmineral supplements lack public myopathy confirmed that statin treatment reduced circulating health recommendations and, except for soy, are not used levels of CoQ10 and that supplementation can raise circulating extensively in the United States, clinicians should keep abreast of levels of CoQ10. 268 However, overall data on CoQ10 blood levels the changing literature on their use. If they are prescribed, frequent are contradictory. checks of reputable resources should be made for contraindications and adverse event reports.

Soy and Soy Isoflavones

Approximately 30% of women use acupuncture, natural estrogens, herbal supplements, or plant estrogens to treat symptoms and discomforts related to menopause. ²⁵² This number was actually higher for women participating in the Study of Women's Health Across the Nation, in which 66% and 37% of the women, respectively, used nutritional remedies or herbal remedies for menopausal symptoms. ²⁵³ Alternative native treatments include phytoestrogens (from soy and red clover), black cohosh, and progesterone creams; only soy and soy isoflavones are discussed here.

Soy protein and soy isoflavones have not been shown to improve vasomotor symptoms of menopause because they do not have sufficient estrogenic activity to have an important impact on vasomotor symptoms of estrogen deficiency in perimenopausal women. ²⁵⁴ With regard to cardiovascular disease prevention, diets substituted with soy protein instead of animal protein can produce significant reductions in LDL-C and triglycerides. ²⁵⁵ Whether this reflects a unique benefit of soy or isoflavones in particular or merely is the result of a reduction in dietary animal protein and fat is unclear. Much of the most favorable evidence on this intervention is observational. A meta-analysis of 10 studies of the effect of soy isoflavones on cholesterol concentrations in wellcontrolled trials substituting soy proteins for dairy or animal protein for at least 14 days found that consumption of soy isoflavones was not related to changes in LDL-C or HDL-C. 256 A review of the benefits of soy protein and isoflavones has been published by the American Heart Association Nutrition Committee. ²⁵⁴ A total of 22 randomized trials evaluated the effects of soy protein and, in aggregate, found a small decrease in LDL-C with no effects on other lipid fractions or blood pressure. Soy isoflavones, also believed to be effective in reducing cardiovascular subclass of polyphenols, are abundant in raw cocoa. disease risk, were evaluated in 19 studies and did not alter lipids or Epidemiologic studies in the Kuna Indians, a population with blood pressure. The American Heart Association concluded that no low cardiovascular event rates compared with their Panamanian meaningful benefit of soy consumption could be demonstrated for neighbors, peaked interest in the antihypertensive effects of HDL-C, triglycerides, or lipoprotein(a). However, the National Cho cocoa. ²⁷¹ In the Zutphen Elderly Study, the median cocoa intake lesterol Education Program Adult Treatment Panel III recommends among users was 2.11 g/day, which was found to be inversely replacing products containing animal fats with soy protein to associated with blood pressure and 15-year cardiovascular and enhance cholesterol lowering.

isoflavones is not warranted and should not be encouraged for synthesis, and promote endothelium-dependent relaxation. cardiovascular disease risk reduction. However, increasing soy Cocoa flavanols enhance flow-mediated dilation through a nitric food consumption in the diet is safe and is encouraged by some oxide-dependent mechanism. 273 Large interindividual variations authorities.

Coenzyme Q10

Coenzyme O10 (CoO10) is involved in cellular oxidative phosphorylation and the generation of ATP. In addition, CoQ10 acts as a free radical scavenger and a membrane

stabilizer. CoQ10 may improve the efficiency of energy 285 production in heart tissue and thus assist the heart during times of physical or oxidative stress. ²⁵⁸⁻²⁶¹ Several meta analyzes reviewing

systematic review on the role of CoQ10 in statin-associated

The largest trial of 1049 patients noted reductions in plasma CoQ10 levels after 52 weeks of treatment with either atorvastatin reduction) or lovastatin (27% reduction). Supplementation with CoQ10 (100 mg/day for 30 days) I decreased muscle pain by 40% in patients with myopathic symptoms associated with statin treatment. However, 200 mg/day in another study did not improve statin-induced ¹⁷ myalgia. ²⁶⁹ Additional well-designed clinical trials are required to address this issue. At this time, routine use of CoQ10 is questionable in statin-treated patients, although it may be an alternative for patients taking statins with myalgia who cannot be satisfactorily treated with other

In general, CoQ10 appears to be safe. No significant side effects have been found, even in studies that lasted a year. CoQ10 chemically resembles vitamin K. However, because vitamin K counters the anticoagulant effect of warfarin, it is not surprising that case reports associate CoQ10 therapy with decreased INR in patients receiving warfarin therapy. 121 On the other hand, 100 mg of CoQ10 daily did not alter the INR in patients taking warfarin in a randomized, double-blind, placebo-controlled, crossover trial. ²⁷⁰ Typical doses range from 100 to 200 mg, two or three times daily. Caution is advised if patients take CoQ10 and warfarin as CoQ10 has the potential to decrease the effectiveness of warfarin.

Recommendation. Routine use of CoQ10 currently is questionable in statin-treated patients, although it may be an alternative for patients taking statins with myalgia who cannot be satisfactorily treated with other agents. CoQ10 otherwise is not recommended for the prevention of cardiovascular disease. Caution is advised in patients taking CoQ10 and warfarin.

Cocoa (Theobroma cacao)

Cocoa belongs to a class of natural compounds, called polyphenols, rich in a number of plant foods. Flavanols, a all-cause mortality. 272 Chocolate and cocoa products inhibit LDL Recommendation. Supplementation with soy or its constituent oxidation and platelet activation, positively affect eicosanoid in absorption of flavanols have been observed, thus limiting interpretation of study results.

> Consumption of flavanol-rich cocoa for 6 weeks in 101 healthy volunteers failed to improve blood pressure (as

shown in earlier studies in normotensive subjects), 274-276 lipid may increase plasma levels of nifedipine. 104 parameters, or C-reactive protein levels. 277 However, chronic administration of 6.3 g/day of dark chocolate containing 30 mg of standardized pharmacological therapies for peripheral vascular polyphenols or matching polyphenol-free chocolate in 44 healthy disease but may be an alternative for patients who do not tolerate prehypertensive or stage 1 hypertensive adults for 18 weeks or are resistant to standard therapy. demonstrated small but significant reductions in systolic and diastolic blood pressures. Reductions in oxidative stress and Horse Chestnut (Aesculus hippocastanum) improved formation of nitric oxide were also observed. 278 A 2week intervention trial administering a flavanol-rich cocoa drink (150 mL twice a day, approximately 900 mg flavanols per day) did not reduce blood pressure or improve insulin resistance in 20 subjects with essential hypertension but did enhance insulinmediated vasodilation. 279

A recent meta-analysis of five randomized trials (173 normotensive and hypertensive subjects) evaluated cocoa intake and blood pressure in studies with a median duration of 2 weeks. Of the five studies, four reported reductions in systolic and diastolic blood pressure with cocoa consumption. The polyphenol content administered was 500 mg/day in three of the five studies and roughly half that in the other two studies. Compared with the cocoa-free control, the pooled decrease was - 4.7 mm Hg in systolic 17blood pressure and - 2.8 mm Hg in diastolic blood pressure for those randomized to cocoa. ²⁸⁰ Similar blood pressure reductions were confirmed in a subsequent meta-analysis. ²⁸¹ Chocolate or cocoa also increased flow-mediated dilation by 4% after acute administration (six studies) and by 1.45% after chronic administration (two studies). 281

Clinical data on consumption of cocoa for cardiac protection appear favorable for blood pressure lowering in individuals with mild hypertension, but the response is not robust. Additional studies are warranted before clinical endorsements can be issued regarding supplemental use beyond incorporation of traditional foods containing cocoa or chocolate into a daily diet plan.

Recommendation. Intake of supplements containing cocoa or its active constituents should be discouraged for disease prevention or reduction of risk factors.

Ginkgo biloba (Ginkgo Leaf Extract)

Ginkgo is one of the most popular herbal therapies in Germany and France, where physicians prescribe it for memory lapses, dizziness, and antiplatelet drugs. anxiety, headaches, tinnitus, and other problems. It has been used for relief of intermittent claudication in patients with peripheral French Pine Bark (Pinus pinaster extract) arterial occlusive disease. 159 Compared with placebo, Ginkgo biloba extract appears to be effective in patients with intermittent claudication (Fontaine stage II peripheral arterial disease). Doses used in clinical trials ranged from 120 to 160 mg/day. Pain-free walking distance and maximal walking distance were often the monitored outcomes of interest. As with other phytomedicines, several constituents of ginkgo extracts may contribute to its therapeutic effect. Ginkgo leaf and its extracts contain several bioactive constituents including flavonoids, terpenoids, and organic acids. ²⁸² Flavonoids reduce capillary permeability and fragility and serve as free radical scavengers. The terpenes, which include the active principle ginkgolides, inhibit platelet-activating factor, decrease vascular resistance, and improve circulatory flow.

A small benefit for ginkgo in the treatment of peripheral arterial disease was confirmed by two meta-analyses reporting a statistically significant increase in walking distance averaging nearly 25 meters. ^{284,285} The most common dosage is 40 mg of standardized extract of ginkgo leaf three times daily. Ginkgo is considered relatively safe and well tolerated and has only a few documented adverse effects, including mild gastrointestinal upset, nausea, dyspepsia, and headache. Ginkgo has been reported to increase both spontaneous and trauma-related bleeding during surgery and other procedures. ^{286,287} Ginkgo does not appear to interact or adversely affect concomitant therapy with cardiac glycosides. The combined use of ginkgo with herbs or drugs with anticoagulant or antiplatelet potential should be avoided. Ginkgo

Recommendation. Ginkgo should not be used in place of

The horse chestnut tree is found worldwide. Horse chestnut seed extract (HCSE) contains saponins, coumarins, flavo noids, and tannins. ²⁸³ Biological activity has been shown to be related to the saponins, notably aescin, which has mild anti- inflammatory properties. ²⁸⁸ The mechanism of action of HCSE involves sensitization to calcium ions to increase the resistance of smallvessel permeability to water. 289

With respect to cardiovascular therapy, research has focused on HCSE for the relief of chronic venous insufficiency . A systematic review of 17 randomized placebo-controlled trials was completed recently. ²⁹⁰ Most studies used a prepared extract containing a daily dose of 100 mg aescin. Leg pain in six trials (n = 543) was significantly reduced compared to placebo. Four of five trials (n = 420) reported a significant decrease in edema in patients treated with HCSE compared with placebo. Calf and ankle circumference were also reduced by HCSE. One trial found HCSE to be as effective as compression stockings. Symptoms of fatigue and tenderness were also reduced with HCSE. An effective dose of HCSE is standardized to 100 to 150 mg aescin daily, which is reduced to 35 to 70 mg daily after improvement. ²⁹¹ Side effects are uncommon but can include gastrointestinal irritation, dizziness, nausea, headache, and pruritus. Contraindications include hypersensitivity to aescin or horse chestnut and renal or hepatic dysfunction. Aescin has antithrombotic effects and might increase the risk of bleeding or bruising. 292 The German Commission E has approved the use of HCSE in chronic venous insufficiency and has listed no precautions or known drug reactions. ²⁸³ Commercial preparations are available in the United States.

Recommendation. HCSE may be an effective alternative to standardized pharmacological therapies for patients with chronic venous insufficiency who are unable to tolerate or are resistant to standard therapy. Monitor with concomitant use of anticoagulants

A common extract made from the bark of the French maritime pine tree is used for chronic venous insufficiency. The strongest evidence for the efficacy of this extract relates to improvement in heart health and treatment of chronic venous insufficiency, but formulations have also been evaluated for the reduction of venous complications of diabetes. 293 The extract consists of a mixture of bioflavonoids and has been shown to have potent antioxidant properties. In a small clinical study of healthy young men, 180 mg/day of French pine bark administered for 2 weeks was shown to augment endothelium -dependent vasodilation by increasing nitric oxide production. 294 In chronic venous insufficiency, it is thought that procyanidins in French tree bark reduce capillary permeability ability by cross-linking capillary wall proteins to reduce edema and microbleeding. 121 The extract also might lower capillary permeability because several of its constituents have antioxidant effects. 121 French tree bark is generally well tolerated, and serious adverse effects have not been recorded. Possible side effects include mild gastrointestinal complaints

and decreased blood glucose levels. It may significantly decrease UNPROVEN TREATMENTS serum thromboxane concentrations, so caution is advised with anticoagulant or antiplatelet agents. 104 An ongoing US clinical trial Bioidentical Hormones will provide additional data on the efficacy of this preparation.

Recommendation. French tree bark extract should not be used as a substitute for standardized pharmacological therapies for suggested that synthetic estrogens plus progestin may pose venous insufficiency but may be an efficacious alternative for patients who cannot tolerate or are resistant to standard therapy.

Omega-3 Fatty Acids

The extensive data from experimental studies, prospective observational studies, and randomized controlled trials are remarkably consistent for a cardioprotective role for omega-3 fatty acids. The anti-inflammatory effects of omega-3 fats may partially explain their cardioprotective benefit because incorporation of these fats in cell membranes promotes vasodilatation, provides antiarrhythmic effects, and promotes vascular patency. 295, 296

Numerous large diet studies support cardiovascular benefits of omega-3 fats. Several meta-analyses have shown a favorable effect of fish and omega-3 fatty acid intake on stroke and fatal coronary heart disease. 297-299 A new pooled analysis of prospective cohort studies that included only healthy individuals without established coronary disease determined that an intake of 250 mg/day of eicosapentaenoic acid (EPA) and docosahexaenoic acid (DHA) reduced risk for coronary heart disease. 300 In addition, a metaanalysis of prospective US studies determined that an intake of approximately 500 mg/day afforded significant protection from coronary disease, 301 which is consistent with the American Heart Association recommendation to consume two fish meals per week for cardiovascular disease protection. ²⁰⁵

The Japan EPA Lipid Intervention Study (JELIS), which was in part a primary prevention trial, randomized more than 18,000 Homeopathy patients with hyperlipidemia on statin therapy to receive 1.8 g/day EPA or control. After 4.6 years, the primary endpoint of "major coronary events" was reduced by 19% in the EPA group. 302 By far the largest secondary prevention trial, the GISSI-Prevenzione medical schools and more than 140 homeopathic hospitals. 322 Study, enrolled more than 11,000 patients who had survived a recent heart attack. Treatment with \$50 mg EPA and DHA per day and 300 mg vitamin E per day resulted in a 20% reduction in total mortality, predominantly by reducing sudden cardiac death (45%).

randomized trials also documented favorable effects on blood pressure, resting heart rate, heart rate variability, and triglyceride Homeopathy has been used worldwide as therapy for an array of levels. 304 However, omega-3 fatty acids compared with placebo did not mitigate the effects of intermittent claudication in a review of six randomized controlled trials (n = 313). This study found some hematological benefits but no improvement in ankle-brachial pressure index or walking distance. 305 Lacking from the evidence base are primary prevention outcome studies in US population groups.

coagulation status. Clinical evidence suggests that fish oil does not affect coagulation even when it is taken with anticoagulant therapy, but diets containing salmon oil, mackerel oil, or cod liver oil have standardized and conventional therapy for the treatment or been reported to prolong bleeding times. 104 Concomitant use of prevention of cardiovascular disease. omega-3 preparations with anticoagulant or antiplatelet drugs should be monitored.

Recommendation. An intake of 1 g/day EPA and DHA for secondary prevention, treatment after myocardial infarction, and Obesity is one of the major risk factors for cardiovascular disease prevention of sudden cardiac death is prudent; higher intakes are and predisposes to numerous cardiac conditions, such as coronary recommended for lowering triglyceride levels. 306 EPA and DHA therapy should be administered under a physician 's care. 307

Results from the Heart and Estrogen/Progestin Replacement Study (HERS) and the Estrogen Replacement for Atherosclerosis trial adverse cardiovascular disease risk to women. 308,309 The Women's Health Initiative trial confirmed that risks were higher in women using conventional hormone replacement therapy and outweighed the benefits for some. 310 Since the Women's Health Initiative, there has been increased interest in the use of bioidentical hormones, which are a derivative of plant extracts chemically modified to be structurally identical to human endogenous hormones.

Many bioidentical hormone products require a prescription because they are compounded by pharmacists. Com pounded preparations of bioidentical hormones may include estriol, estrone, estradiol, testosterone, DHEA, thyroxine, cor tisol, or progesterone. Some clinical studies and studies in postmenopausal cynomolgus monkeys suggest beneficial physiological changes in lipid 311-313 blood clotting, 312,314,315 and coronary metabolism, vasoreactivity. 316-318 However, there is minimal scientific evidence, and rigorous controlled clinical studies are lacking. The bioidentical hormones can be expected to have the same adverse effect profile as that of conventional hormone replacement therapy. ³¹⁹ There are proponents who believe that bioidentical hormones remain the preferred method of hormone replacement therapy until it is proven otherwise. 320,321

Recommendation. Bioidentical hormones are not recommended for the reduction of risk in patients with cardiovascular disease.

Homeopathy is a distinct formal system of medicine that has a long history of use in America. At the turn of the twentieth century, 8% of US physicians used homeopathy. There were 20 homeopathic Samuel Hahnemann introduced the practice based on the principle of similars. That is, if a substance in large amounts caused a certain condition, then the same substance in small amounts could cure the condition. Treatments are usually patient specific individualized. Most homeopathic products contain little or no Short-term administration of 1 to 2 g/day of EPA and DHA in active ingredient and therefore possess little risk and require minimal oversight by the Food and Drug Administration. chronic and acute cardiovascular conditions. However, data from randomized controlled trials and meta-analyses have major methodological limitations, and therefore interpretation of results of therapeutic efficacy is questionable. 323 Five systematic reviews and meta-analyses evaluated clinical trials of the effectiveness of homeopathic remedies compared with placebo. "The reviews found that, overall, the quality of clinical research in homeopathy It is unclear to what extent omega-3 fatty acids affect is low. But, when high-quality studies were selected for analysis, a surprising number showed positive results." 324

Recommendation. Homeopathy should not be used in place of

Supplements for Weight Loss

disease, heart failure, and sudden death. No prospective weight loss studies have demonstrated increased survival. However, strong evidence does suggest that weight loss in overweight and obese individuals reduces the risk for

288 development of diabetes and cardiovascular disease. 325 A higher body mass index has been associated with a higher likelihood of angiographic progression of coronary artery disease. 326

Liquid protein diets and very-low-calorie diets have been associated with adverse cardiac events independent of the cardiovascular diseases. 327

Survey data suggest that adults with higher body mass index are no more likely to use CAM therapies for weight loss than normal-weight individuals are. 328 Herbal antiobesity products as **CONCLUSION** a class are heterogeneous in composition and often have unpredictable levels of active ingredients. Potentially harmful are Integrative medicine consists of a large number of diverse practices would be consumed ¹⁷ if the ingredients were taken individually, trials showing efficacy are relatively scarce (Table 17-4). presumably to increase energy expenditure. These include ingredients such as guarana, kola nut, cocoa, and yerba mate. The

cardiovascular and central nervous system effects of ephedra are magnified when it is consumed with caffeine-containing products. ¹⁸⁵ Last, there is also the potential for intentional as well as unintentional (misidentification) adulteration of ingredients, including the addition of drugs or other natural products. 329

Recommendation. The mainstay for weight loss remains biological and nutritional value of the products' constituents. decreased calorie intake and increased physical activity or caloric 325 Current prescription weight loss products, such as sibutra expenditure. The only appropriate supplements for weight loss are mine hydrochloride and orlistat, have shown some success those prescribed by knowledgeable clinicians, which may include a but may be contraindicated in some patients with multivitamin-mineral supplement to compensate for those nutrients lost in calorie-restricted diets.

toxic herbs such as Aristolochia species, ephedra or ma huang and used widely by adults and less so by children in the United States bitter orange (Citrus aurantium) because of their sympathomimetic today. Interest in these non-traditional therapies stems from general stimulatory activity, and herbal laxatives (anthraquinones, dissatisfaction with current Western methods of healing and anthrones, and dianthrones) that may cause hepatotoxicity and increased ownership that the public has assumed in their own health nephrotoxicity or electro lyte depletion. A number of herbal care. Furthermore, the American public is bombarded with weight loss blends contain caffeine in higher levels than typically advertisements and claims of efficacy, when in reality strong clinical

	Blood Pressure	Blood Lipids	Stress Reduction	Coronary Artery Disease or Angina Pectoris	Venous or Vascular Insufficiency	Obesit
Traditional Chinese medicine Acupuncture, acupressure, moxibustion	Yes	Yes	Possibly	Yes		Yes
Energy therapies Tai Chi, Qigong Chinese herbs	Yes		Possibly	Yes		
Danshen		Yes* Yes*		Yes* Yes*		
Compounded <i>Salvia</i> Suxiao jiuxin wan Tongxinluo Red yeast rice		Yes			No No	
Ayurvedic medicine	V		Deseibb			
Yoga Herbs	Yes		Possibly			
Garlic	Possibly	No				
Gu qqui	1 0331019	No				
99 Terminalia arjuna		No				
Meditation and stress reduction	Yes		Yes			
Naturopathic medicine	Lacks evide	nce-based da	ata to determine eff	ïcacy		
Chelation therapy				No		
Nutritional supplements						
B vitamins				No		
Antioxidants				No		
Vitamin D				No		
Magnesium	No			No		
Omega-3 fatty acids	Yes	Yes		Yes		
Nonvitamin, nonmineral supplements		NI.		NI.		
Soy		No		No		
Coenzyme Q10 Cocoa	No			Yes*		
Ginkgo biloba					Yes*	
Horse chestnut seed extract French pine bark					Yes Yes	
Bioidentical hormones		No		No		
	Lacks evide	nce-based da	ata to determine eff	ïcacv		
Homeopathy	Edono ovido					

^{*}Indicates beneficial use in some patients under some conditions; review contraindications and concomitant medications

nutritional approaches, the mechanisms of action are beginning to be elucidated. This fundamental approach to demonstrate how these therapies work along with rigorous clinical trials will be necessary 24. Petrovic P, Kalso E, Petersson KM, Ingvar M: Placebo and opioid analgesia—imaging a shared before the medical community accepts these new therapies as advantageous over current Western approaches to medical practice. 25. Pariente J, White P, Frackowiak RS, Lewith G: Expectancy and belief modulate the neuronal However, despite the lack of basic and clinical information, there 26. may be utility in adopting an integrative approach to health care that employs both conventional and nonconventional treatments, if they 27. have a positive impact on chronic conditions for which Western medicine often has proven to be inadequate.

There is emerging evidence that many of the therapies can reduce stress and associated cardiovascular responses, such hypertension, that frequently accompany disease. Further more, the 30. preventive approaches advocated in naturopathic, traditional Chinese, and Ayurvedic medicine seem to coincide with Western epidemiologic studies indicating that reduction of cardiovascular 32. Longhurst JC: Integrative cardiology: mechanisms of cardiovascular action of acupuncture. In risk by improving nutrition, increasing exercise, and lowering body weight can reduce future morbidity. In fact, some of the most promising data in the areas of nutrition and positive health outcomes relate to dietary patterns and not dietary supplements. There are insufficient data to justify an alteration in public health 34. Zhou W, Hsiao I, Lin V, Longhurst J: Modulation of cardiovascular excitatory responses in rats by policy from one that emphasizes food and diet to one that emphasizes supplements. 330 However, use of well-characterized dietary supplements in targeted individuals at risk for development of cardiovascular disease may serve to complement conventional 36. therapies. We believe that to foster healthy lifestyles in their patients, clinicians will need to become more involved in helping patients remove barriers and adopt lifestyle measures that meet their needs 38. and at the same time reduce their cardiovascular risk.

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APPENDIX to Chapter 17

Reported Uses and Potential Interactions Between Dietary Supplement Products and Conventional Medications

Supplement Arginine	Reported Uses Improve peripheral and coronary blood flow and as adjuvant therapy for CAD, CHF, PVD, angina pectoris, and hypertension	Active Ingredient L -Arginine	Potential Interactions Has been shown to have additive hypotensive effects if combined with hypertensive drugs, but data is limited	Comments and Suggested Guidelines of Use L-Arginine is the precursor of nitric oxide and has been shown to improve coronary and brachial artery endothelial function. Short-term efficacy for intermittent claudication due to PVD was not replicated in a longer term trial. Efficacy of oral L-arginine for CHD awaits confirmation.
Artichoke leaf extract (Cynara cardunculus)	Lower blood cholesterol	Phenolic acids (chlorogenic acid, cynarin, and caffeic acid), sesquiterpene lactones, and flavonoids (scolymoside, cynaroside, and luteolin)	No known interactions with herbs and other dietary supplements, drugs, foods, or laboratory tests	In the United States, artichoke has GRAS use in foods. Few data from rigorous clinical trials exist. Beneficial effects are reported but not compelling. Cynaroside and its derivative, luteolin, might also indirectly inhibit HMG-CoA reductase. Potential adverse effects include flatulence and allergic dermatitis.
Astragalus, root (Astragalus membranaceus)	Used as an anti- inflammatory, antioxidant, diuretic, vasodilator, and hypotensive agent	flavonoids (isoflavones,	No known interactions with herbs and other dietary supplements, drugs, foods, or laboratory tests. However, certain isoflavones affect alcohol dehydrogenase.	Astragalus is most commonly used in combination with other herbs. Small clinical studies have found astragalus to be effective for relief of angina pectoris. As astragalus may stimulate immune function, theoretically it might decrease the effects of immunosuppressive therapy.
Carnitine	Reduces symptoms of angina pectoris and PVD Adjunct therapy for heart failure	L -Carnitine, acetyl- L -carnitine, and propionyl- L -carnitine	I Concomitant use with acenocoumarol may potentiate (decrease) anticoagulant effects. It is unknown if interactions occur with warfarin.	L-Carnitine is naturally found in the body. D-or DL-carnitine should not be substituted for L-carnitine. Potential side effects include GI upset, diarrhea, body odor, and seizures. The FDA approved L-carnitine for use in primary carnitine deficiency. CHF clinical trials are underway in Italy. Monitor INRs in patients receiving anticoagulant therapies. ACC/AHA Practice Guidelines note that the effectiveness of propionyl-L-carnitine as a therapy to improve walking distance in patients with intermittent claudication is not well established (Level of Evidence: B).



				Comments and Suggested Guidelines of
Supplement Ginseng root, Asian (Panax ginseng)	Reported Uses Acts as an "adaptogen" to enhance energy levels and to relieve stress and eases symptoms of anxiety Antihypertensive Improves control of blood glucose	Active Ingredient Ginsenosides and triterpenoid saponins are the main active components derived from the root	Potential Interactions Interacts with P-450 enzyme system Potential to interact with anticoagulant and antiplatelet drugs and corticosteroids, calcium channel blockers, digoxin, and methyldopa Interacts with MAO inhibitors, stimulants, and phenelzine sulfate (an antidepressant) May enhance the effect of hypoglycemics	Use
Green tea (Camellia sinensis)	Lower blood cholesterol Lower blood pressure Improve cognitive performance Weight loss	Parts used include leaf bud, leaf, and stem Polyphenols (flavanols, such as catechins, flavandiols, flavonoids) and phenolic acids and caffeine	Green tea contains catechins and caffeine as well as vitamin K and thus may interfere with anticoagulant or antiplatelet drugs and antagonize the effect of warfarin. Verapamil has been reported to T plasma caffeine concentrations by 25%.	A green tea health claim for CVD was recently rejected by the FDA. Patients receiving warfarin need to be routinely questioned about their intake of vitamin K-containing foods, beverages, and dietary supplements. Monitor INR in patients who habitually consume green tea and are taking warfarin.
Hawthorn (Hawthorn)	Reduces symptoms of mild CHF, ischemic heart disease; reduces the risk of arrhythmias and atherosclerosis	Hawthorn leaf extract (leaves, fruit, and flowers) contains procyanidins, flavonoids, triterpenoids, catechins, and aromatic carboxylic acids.	Has been hypothesized to potentiate the effects of digitalis and have additive effects when combined with beta blockers, calcium channel blockers, and nitrates to T vasodilation	In Germany, hawthorn is prescribed for "mild cardiac insufficiency." It is generally well tolerated and safe for short-term use. Most common side effects are GI upset, fatigue, and dizziness. Recently noted increased risk of death and hospitalization in NYHA stage II or III heart failure patients taking 450 mg twice daily participating in a long-term clinical trial
Horse chestnut seed extracts (Aesculus hippocastanum)	Reduces swelling and discomfort due to chronic venous insufficiency, varicose veins and phlebitis	Aescin, a triterpene saponin, and aesculin, a glycoside, which is also a hydroxycoumarin	May T effects of anticoagulants and antiplatelet drugs Possibly induces hypoglycemic effects Do not administer with drugs known to cause nephrotoxicity.	Most widely prescribed oral antiedema venous remedy in Germany Treatment for 1 to 3 months may be required before full therapeutic effects are apparent. Side effects are uncommon, but GI irritation and toxic nephropathy may occur. Monitor individuals taking warfarin. Monitor blood glucose levels in diabetics.
Hu zhang (Polygonum cuspidatum)	CVD and lipid lowering	Hu zhang root contains flavonoids, phenolic acids and their derivatives, tannins, stilbenes (resveratrol), and anthraquinones.	Resveratrol has been shown to have antiplatelet effects and therefore could T risk of bleeding. Do not administer with anticoagulant or antiplatelet drugs.	There is limited reliable information available about the effectiveness of Hu zhang. Hu zhang is considered to be one of the richest sources of <i>trans</i> -resveratrol.

Continued

Supplement Jiaogulan	Reported Uses Reduce blood cholesterol and	Active Ingredient Jiaogulan leaves contain triterpene	Potential Interactions Clinical trial data is lacking.	Comments and Suggested Guidelines of Use
(Gynostemma pentaphyllum)	blood pressure and improve coronary and cardiovascular functions	saponins (gypenosides), many of which are similar to ginsenosides found in <i>Panax</i> <i>ginseng</i>	Ü	Jiaogulan grows wild in China, and it is a newcomer to TCM. Most of the evidence regarding the pharmacological effects of jiaogulan comes from preliminary animal or in vitro research. Older studies in humans suggest cholesterol-lowering effects. Newer clinical data suggest an antidiabetic effect.
Kudzu (Pueraria Iobata)	Lower blood pressure Reduces symptoms of angina pectoris	Kudzu (root, flower, and leaf) contains isoflavones daidzin, daidzein, puerarin, genistin, and genistein	No side effects have been reported in clinical studies. Theoretically, concomitant use with anticoagulant or antiplatelet herbs or drugs might increase the risk of bleeding. Dose-dependent Antabuse like activity with ethanol.	Kudzu has been used medically in Chinese medicine since 200 Bc . Kudzu was brought to the southeastern United States in the late 1800s to help prevent soil erosion but has now turned into one of the most invasive species in this region. The kudzu extract puerarin has been used intravenously to treat ischemic stroke.
Lycium (Lycium chinense)	Improve circulation and lower blood pressure and blood glucose levels	Lycium (dried berries and root bark) contains beta-sitosterol; the bark also contains kukoamine, which may lower cholesterol; also contains beta carotene, niacin, pyridoxine, and ascorbic acid	Clinical data is lacking.	Lycium is a native Chinese deciduous shrub with bright red berries that has been promoted to increase longevity. Lycium is usually taken orally as a tea. Two case reports suggest interaction with warfarin therapy.
Policosanol	Lower blood cholesterol For peripheral vascular insufficiency and intermittent claudication	Derived from a variety of plant sources and contains a mixture of waxy alcohols	Policosanol may inhibit platelet aggregation, so use caution with anticoagulant and antiplatelet drugs or supplements such as garlic and ginkgo and high doses of vitamin E.	Research for hypercholesterolemia is inconsistent and contradictory. Policosanol appears to be well tolerated; it may cause erythema, migraines, insomnia, irritability, dizziness, upset stomach, and rash. INR monitor in patients taking warfarin. Discontinue use before surgery.
Schisandra (Schisandra chinensis)	Reduces blood pressure and elevated cholesterol	Active constituents in the fruit include lignans (schizandrins, schizandrols, gomisins, schizandrers, schisantherins, wuweizisu) and citral, stigmasterol, and vitamins C and E	Induces cytochrome P-450 2C9 (CYP2C9), possibly altering metabolism of fluvastatin, irbesartan, losartan, and warfarin	It may cause heartburn, acid indigestion, decreased appetite, stomach pain, allergic rashes, and urticaria in some patients.
Selenium	Prevent oxidative stress	Selenomethionine is generally considered to be the best absorbed and used form of selenium	Theoretically, combining selenium with anticoagulant or antiplatelet drugs might T INR and bleeding times through multiple pathways.	Selenium appears to be safe when it is taken short term in amounts below the upper intake level of 400 ^ g/day. Selenium toxicity can elevate the ST segment and cause T wave changes on EKG characteristic of myocardial infarction. Long-term use has been shown to increase the risk of developing type 2 diabetes and to increase mortality. AHA notes that "antioxidant vitamin supplements or other supplements such as selenium to prevent CVD are not recommended."



Reported Uses Supplement Prevention of CVD (CHD, Vitamin E stroke)

Active Ingredient Family of 8 fat-soluble antioxidants, Anticoagulant or antiplatelet drugs a -tocopherol most active form

Potential Interactions and herbal medicines with anticoagulant and antiplatelet potential have 1 INR.

High doses of vitamin E (> 800 units/day) can antagonize the effects of vitamin K and 1 the risk of bleeding.

Comments and Suggested Guidelines of

Relatively nontoxic; safe when used orally in amounts not exceeding the UL of 1000 ma/dav

Conversion factor: 1 mg = 1.26 IU;

RDA of 15 mg = 22 IU (natural form) or 33 IU (synthetic form) Possible T risk of bleeding for those

taking anticoagulants or with vitamin K deficiency

Possible T risk of hemorrhage in high-risk aroups

Monitor INR in patients taking warfarin and vitamin E > 800 IU daily

AHA notes that "antioxidant vitamin supplements or other supplements such as vitamin E to prevent CVD are not recommended"

ACC/AHA, American College of Cardiology/American Heart Association; CAD, coronary artery disease; CHD, coronary heart disease; CHF, congestive heart failure; CVD, cardiovascular disease; EKG, electrocardiogram; FDA, Food and Drug Administration; GI, gastrointestinal; GRAS, generally recognized as safe; HMG-CoA, hydroxymethylglutaryl-coenzyme A reductase; INR, international normalized ratio; MAO, monoamine oxidase; NYHA, New York Heart Association; PVD, peripheral vascular disease; RDA, recommended daily allowance; TCM, traditional Chinese medicine; UL, upper tolerable limit.

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KEY POINTS

- Most large prospective cohort studies show that regular, moderate consumers of alcohol have considerably lower risk for development of coronary heart disease and other types of CVD than do abstainers.
- Numerous mechanisms have been demonstrated in basic scientific studies and limited human trials to explain the effects of both alcohol and the polyphenols in some alcoholic beverages on reduction of CVD risk; these include beneficial effects on lipids, coagulation, fibrinolysis, inflammation, glucose metabolism, and endothelial function.
- Evidence is accumulating that moderate drinkers are also at lower risk of other chronic diseases (including diabetes, dementia, and osteoporosis) and have lower all-cause mortality rates than those of nondrinkers.
- In contrast to potential health benefits from moderate drinking, there are many adverse health and societal effects of the abuse of alcohol. Advice to patients regarding alcohol consumption will vary according to their individual characteristics (age, sex, risk factors) but should always be based on scientifically sound and balanced data.
- Any recommendations regarding moderate alcohol consumption for the prevention of CVD for individuals should be based on consultation with the health care provider.

CHAPTER 18

Effects of Alcohol on Cardiovascular Disease Risk

R. Curtis Ellison

EPIDEMIOLOGICAL EVIDENCE RELATING ALCOHOL TO CORONARY HEART DISEASE

There are notable exceptions to the inverse association between alcohol and coronary artery disease occurrence or mortality in epidemiological studies. As would be expected, longitudinal studies based primarily on young people tend to have few instances of CVD and may show no protection against CHD.¹² ¹³ ¹⁴Studies from eastern Europe, where very heavy drinking is the norm, often do not show lower rates of CHD among drinkers. In some studies in which binge drinkers are combined with regular moderate drinkers, lower rates for "moderate " drinking have not been seen. 13,14 Moreover, if the range considered to be moderate drinking extends to more than three or four typical drinks per day, an inverse association may not be seen among drinkers. The meta-analysis by Corrao and associates 15demonstrated that essentially all studies show lower CHD for consumers of up to 12.5 g of alcohol per day, but some fail to show lower rates when consumption of up to 25 or 50 g/day is considered the moderate

As described by Wannamethee and Shaper, ¹⁶it is essential that ex-drinkers not be included with lifetime abstainers in the nondrinking category, as the former tend to have higher rates of many types of disease that may falsely increase a putative beneficial effect among

¹²g of alcohol, the average amount in a "typical drink"). The relative risk crossed

^{1.0 (}the risk for nondrinkers) between 72 and 89 g of alcohol per day (Fig. 18-1). Starting in the early 1970s, many scientists began to publish prospective data

A lower risk for CHD and other cardio vascular disease (CVD) among moderate drinkers has been shown in populations with a low intake of alcohol, such as the Chinese, 6 * * and in those with typically high alcohol intake, as in Scandinavia. 7.8 Data from the latter show that heavy drinking is associated with increases in sudden death. 9 Rimm 10 stated that the evidence to support the hypothesis that the inverse association of alcohol to CHD is causal, and not confounded by healthy lifestyle behaviors, is very strong. This evidence includes the fact that moderate alcohol consumption reduces CHD and mortality in individuals in diverse cultures with varying drinking habits; among subjects with hypertension, diabetes, and existing CHD;

moderate drinkers. Poikolainen ¹⁷has pointed out the importance of identifying abusive drinkers in a population before attempting to evaluate the association between moderate drinking and CVD. As will be described later, some studies suggest that African Americans may not show a lower risk of CVD from moderate drinking.

LACK OF RANDOMIZED CLINICAL TRIALS OF ALCOHOL CONSUMPTION AND CORONARY HEART DISEASE EVENTS

In lieu of randomized clinical trials of alcohol intake and cardiovascular outcomes , physicians are forced to rely primarily on associations shown in observational

and among otherwise healthy individuals. 10 Mukamal and coworkers 11 demonstrated among "very healthy" subjects (who did not smoke, exercised, ate a good diet, and were not obese) that those who drank moderately had a much lower relative risk of CHD (0.38; 95% CI, 0.16-0.89) than that of similar subjects who did not consume any alcohol.

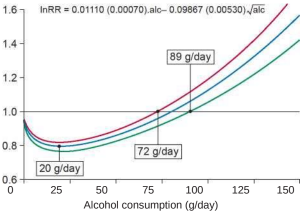


FIGURE 18-1 Association between alcohol intake and CHD: a meta-analysis. Functions (and corresponding 95% CIs) describing the dose-response relationship between alcohol consumption and the relative risk of CHD obtained by pooling from a meta-analysis the 28 cohort studies at

which a > 15 quality score was assigned. The fitted model (with standard errors in parentheses) and three critical exposure levels (nadir point, maximum dose showing statistical evidence of protective effect, and minimum dose showing statistical evidence of harmful effect) are also reported. (From Corrao G, Rubbiati L, Bagnardi V, et al: Alcohol and coronary heart disease: a meta-analysis. Addiction 95:1505, 2000. Reprinted with 18 permission.)

studies, experiments of the effects of alcohol on risk factors, and animal experiments. Also, results among subjects in prospective studies who change their drinking habits during follow-up may be informative. Several studies have demonstrated that when former abstainers begin to consume alcohol, their subsequent risk of CVD decreases. Gronbaek and colleagues, 17 in a prospective study of more than 14,000 subjects, showed that moderate drinkers had a lower risk of total mortality than nondrinkers or heavy drinkers. Furthermore, based on alcohol assessments 5 years apart, they found that in comparison with stable drinkers, subjects who reduced their drinking from light to none increased their mortality risk (RR = 1.40; 95% CI = 1.00-1.95), and those who went from nondrinking to light drinking reduced their risk (RR = 0.71; 95% CI = Whereas most of the original studies relating alcohol intake to CHD 1.10, CI: 0.92, 1.31). ¹⁸

Former abstainers who reported moderate intake in fol low-up examinations in the Atherosclerosis Risk in Communities (ARIC) study subsequently had a 38% lower chance for development of Freiberg and coworkers 35 reported lower total mortality among CVD than did their persistently nondrinking counterparts. 19 In a African American hypertensives who consumed alcohol than large population-based cohort of women in Australia, moderate among those who did not drink, but a similar relationship was not drinkers showed the best self-reported state of overall health, seen among nonhypertensive African Americans. whereas those who stopped drinking during follow-up (even those with no reported comorbidity) showed a decrease in their ratings of overall health ²⁰; as in most studies, however, the reasons that the subjects stopped drinking were not known.

GENDER, AGE, AND ETHNICITY

Gender

Women are known to show decreased tolerance to alcohol, which is thought to relate to lower levels of alcohol-clearing enzymes (especially alcohol dehydrogenase), their generally smaller body size, and the greater proportion of their body consisting of fat, which metabolizes alcohol less well than muscle does. Both the beneficial and adverse effects of alcohol appear at lower dosages for women. 10,21,22 Hence, guidelines generally suggest that moderate drinking for women should be only about half the amount of alcohol for men. ²³ From a meta-analysis, Di Castelnuovo

and coworkers 24 found that women drinkers showed an increase in mortality over that of nondrinkers above about 2 drinks/day, whereas the increase among men was at about 3)2 drinks/day. In the Australia Longitudinal Study on Women's Health, with approximately 12,000 subjects aged 70 years and older, Byles and colleagues 25 found that nondrinkers and women who rarely drink had a significantly higher risk of dying (HR = 1.94; 95% CI, 1.4-2.6) during the survey period than did women who consumed alcoholic beverages at a level of one or two drinks per day; the best general health status and highest levels of physical functioning were among those drinking at this level for 3 to 6 days per week.

Age

Alcohol has been shown to be associated with CHD in the elderly as well as in middle-aged people. In the Cardiovascu lar Health Study, subjects older than 65 years who consumed 14 or more drinks per week had a risk of incident myocardial infarction or cardiac death about 40% lower than that of abstainers. ²⁶ People tend to drink less as they age, ^{27,28} and most current recommendations state that individuals older than 65 years should drink less than what is considered moderate for younger people. ²³ On the other hand, two analyzes comparing the health effects among elderly subjects consuming one or two drinks per day, in comparison with those consuming only one drink per day or less, have not shown higher morbidity or mortality among those consuming more alcohol. 29.30

Tolvanen and colleagues 31 evaluated 10-year mortality according to alcohol intake among subjects initially aged 60 to 99 years, of whom 50% of men and 40% of women died during followup. After adjustment for age, sex, educational level, marital status, chronic diseases, functional ability, and smoking, the relative risk for mortality of frequent drinkers (versus nondrinkers) was 0.6 (95% CI, 0.4, 0.8); for occasional drinkers, 0.7 (0.5, 1.0); and for exdrinkers, 1.1 (0.8, 1.7).

Ethnicity

0.44-1.14). 17 In the Health Professionals Study, with repeated were in Europeans and European Americans, a large number of alcohol assessments, subjects reporting an increase of > 10 g/day in studies have found essentially the same effects among Asians. ^{32,33} alcohol had a decrease in risk of subsequent myocardial infarction Data on African Americans are limited, and Sempos and colleagues (RR = 0.55; CI 0.33-0.91), whereas there was a tendency for a slight ³⁴ found no inverse association between alcohol and CVD risk increase in risk for subjects decreasing their alcohol intake (RK = among African Americans. Klatsky and associates, 32 however, found similar inverse associations between alcohol and heart failure in the Kaiser Permanente cohort among whites, Asians, and African Americans. Using data from the Women's Health Initiative,

IMPORTANCE OF **PATTERN** OF **ALCOHOL CONSUMPTION**

DIFFERENCES IN RESPONSE TO ALCOHOL BY It has become very clear that the amount of alcohol consumed (within limits) is not as important in terms of health effects as is the pattern of consumption. The two key aspects of the pattern are the frequency of drinking and a large number of drinks on a single occasion (binge drinking). Mukamal and coworkers demonstrated among people who had suffered a

myocardial infarction that those who drank moderately without abstainers, and the inverse association between moderate binge drinking had 30% fewer deaths, whereas those who reported drinking and CHD remains, as described by Rimm and Moats 10 binge drinking (defined in this study as consuming three or more and others. 46 drinks within 1 to 2 hours) had death rates even higher than those of teetotalers.

substantial increase in risk. 38

Cross-cultural prospective studies have shown that simply alcohol intake and CVD and all-cause mortality. 51 correlating the average amount of alcohol consumed with CVD outcomes is inadequate. 39 In some cultures, daily drinking is the number of lifestyle factors that were considered to increase the risk norm, but in others, alcohol consumption is generally confined to of CHD; these included hypertension, increased body mass index Friday and Saturday nights; whereas benefits are seen in the (BMI), diabetes, depression, sleep disturbances, smoking, former, they are absent in the latter. 40 Data are mixed on whether physical inactivity, poor life satisfaction, psychological distress, consumption of alcohol in conjunction with food affects the trait anxiety, independent and dependent life events, longer outcome, but elevated risk for hypertension has been reported in length of working hours, low levels of job control, job strain, and one study only for individuals who drink outside of meals. 41 effort-reward imbalance. Most of these conditions were either the Furthermore, Gorelik and associates 42 found in an intervention same between lifetime abstainers and light to moderate drinkers study that red wine served with a high-fat meal inhibited the or more common among drinkers than among abstainers. Thus, postprandial increase in serum and urinary levels of cytotoxic lipid these authors concluded that none of the large number of lifestyle peroxidation products.

IS IT ALCOHOL OR ASSOCIATED LIFESTYLE **FACTORS THAT RESULT IN LOWER RISK FOR CORONARY HEART DISEASE?**

Many lifestyle factors are associated with moderate drinking. In all epidemiological studies, socioeconomic factors are strongly and inversely related to the risk of CHD; better-educated individuals and those with higher income show lower rates of CHD (and most other diseases), presumably because of their more moderate lifestyles. The extent to which residual confounding by such factors may explain the health-protective effects of moderate drinking shown in observational studies has provoked considerable debate.

As a follow-up to concerns from earlier work by Shaper and colleagues, 43 Fillmore and coworkers 44 have described apparent "errors" in prospective studies that call into question their conclusions about an inverse association between alcohol and CHD. One of their arguments is that exposure measures may not be as accurate in prospective cohort studies as in case-control studies, a point not supported by most epidemiologists. In general, there is greater possibility of misclassification of exposure in casecontrol studies than in follow-up studies, as the former may EFFECTS OF ALCOHOL ON SUBJECTS "introduce a number of subtleties and avenues for bias that are ALREADY DIAGNOSED WITH CORONARY absent in typical cohort studies." 45

The choice of the referent group in an epidemiological study is important, and the nondrinker category should not mix ex-heavy Janszky and coworkers 54 reported results from serial drinkers (who usually have higher risks for many diseases) with quantitative coronary angiography studies done among middlelifetime abstainers. In a population in which abstinence is rare, aged choosing very light or occasional drinkers as the referent group may be preferable. A key criticism of

existing observational studies of alcohol and CHD relate 303 to the inclusion in some of ex-drinkers in the nondrinking category. 44 This is indeed an important problem, but many recent reports have limited the referent, nondrinking group to lifetime

An approach for dealing with potential residual confounding by social class is to limit analyzes to subjects who are very similar in Studies of the effects of alcohol intake on CHD, obesity, socioeconomic terms, as has been done for nurses, ⁴⁷ health cognitive function, and other diseases show that more frequent - professionals, ⁴⁸ and business executives. ⁴⁹ In each of these groups drinkers have the best outcomes; for subjects consuming the same of subjects, moderate drinking was associated with lower CHD average amount of alcohol during a week, those consuming alcohol risk. Lee and coworkers 50 evaluated a large variety of measures of every day may have up to 50% lower risk of disease than those lifestyle in seeking to determine if the lower total mortality risk in consuming the same amount on only 1 or 2 days. 36 In a meta-moderate drinkers is due to the alcohol itself or to associated analysis based on the only six prospective studies in the literature healthy lifestyle factors. Adjusting for a large number of such that allowed such an analysis of pattern of drinking and CHD, factors and using sophisticated analytical techniques, the authors Bagnardi and coworkers 37 reported that in comparison with not found that moderate drinking (versus not drinking) was still consuming alcohol, regular drinking (even at fairly high levels of associated with a 38% lower mortality risk. 50 Friesema and consumption) was associated with lower risk of CHD; irregular colleagues 51 reviewed these relationships among subjects in a heavy consumption, or binge drinking, increased the risk. ³⁷ In a prospective case-cohort, the Lifestyle and Health Study, consisting population of light to moderate drinkers, alcohol consumption in of 16,210 men and women between 45 and 70 years of age. The general was associated with decreased risk of acute myocardial - authors concluded that the difference in lifestyle between moderate infarction in women, but episodic intoxication was related to a drinkers and 1 both never drinkers and former drinkers was only a partial explanation of the observed inverse relationships between

> Poikolainen and associates 52 evaluated the effects of a large factors evaluated is likely to be the reason that abstainers have higher rates of CHD, which supports the theory that it is the alcohol that is associated with less disease. 52

Rimm ¹⁰ summarized the research data on this topic by stating that results from mechanistic studies provide substantial support for the hypothesis that moderate alcohol intake reduces the risk of CHD and that all beverages containing alcohol have shown beneficial effects. He concluded, "The 'sick-quitter' hypothesis and the concern that moderate drinkers lead a healthier lifestyle may explain a small proportion of the benefit attributed to alcohol in some studies, but recent studies which have removed sick quitters, updated alcohol and covariate information on diet and lifestyle factors, and separately documented benefits of alcohol among healthy and unhealthy populations further add to the evidence that moderate alcohol consumption is causally related to a lower risk of CHD." 10 Sorensen and coworkers 53 stated that because alcohol consumption is so common in most Western cultures, it would be exceedingly difficult to carry out a sufficient number of randomized controlled trials to evaluate adequately the effects of alcohol on CVD outcomes. They pointed out that medical decisions must often be made on the basis of observational studies. 53

HEART DISEASE

- Improvement of blood lipids (marked increase in HDL, slight decrease in LDL)
- Improvement of coagulation, fibrinolysis
- Improvement of endothelial function
- Activation of genes for fibrinolysis, eNOS
- Improvement of ventricular function
- Reduction of inflammation
- Improvement of glucose metabolism
- Adverse effects on blood pressure, especially for heavier drinking

women at 3 to 6 months and at 3 years after an acute myocardial infarction. They found that moderate alcohol consumption (more than 5 g/day) was protective of coronary atherosclerosis 18 progression. Furthermore, restenosis after stenting has been remain in a favorable state all the time. Unfortunately, many found to be lower in patients with CHD who continue to consume alcohol. 55 The risk of complications (including death) after an acute myocardial infarction has been shown to be lower for subjects who unhealthy way to drink. remain drinkers than for those who abstain. 56,57 Such a protective effect seems to be lost when the subject consumes alcohol as binge- Other Mechanisms drinking. 13

BIOLOGICAL MECHANISMS FOR CARDIOVASCULAR EFFECTS

Box 18-1 lists some of the key mechanisms by which moderate alcohol consumption, as well as the intake of polyphenols in red wine and other beverages, may lead to a reduction in the risk of CHD and other types of CVD.

Lipids

 $CVD\ relates\ to\ induced\ changes\ in\ plasma\ lipid\ profile,\ especially\ lesion\ progression\ ^{106,107}\ ;\ inhibition\ of\ endothelin\ 1\ synthesis\ ^{108}\ ;$ increases in high-density lipoprotein cholesterol (HDL-C) and its downregulation of tissue factor gene transcription in cultured subtypes (HDĽ 2, HDĽ 3). 58-61 Numerous observational and human endothelial cells and monocytes 109,110; and inhibition of experimental studies have confirmed a strong, direct association smooth muscle cell proliferation. 111,112 Research studies between alcohol intake and HDL. 8,62,63 Although it was initially increasingly show a strong protective effect from alcohol and believed that only certain HDL subtypes were increased by alcohol polyphenols against various indices of inflammation. 113,114 Wang (and that those had little effect on CVD risk), alcohol intake has and colleagues 115 have shown a U-shaped association between been shown to increase both HDL 2 and HDL 3, and both play a role alcohol and high-sensitivity C-reactive protein (hsCRP) levels; a in CVD prevention. 64-66 In addition, most studies also show slightly proportional odds model analysis showed an odds ratio for lower LDL concentrations, especially small-particle LDL, and non-HDL lipoprotein levels among moderate drinkers. 67-70 Oxidation of 70 g/day (about 1% to almost 6 drinks) in comparison with lipids has been shown to be decreased by alcohol or polyphenols in nondrinkers. Alpert and coworkers 116 found that CRP was lowest many studies 71-75 but not in all. 76

Platelet Aggregation and Thrombosis

As described by Booyse and colleagues, 77 other changes in p38 MAPK signaling by alcohol as well as by certain wine vascular, myocardial, hemostatic, and endothelial cell functions polyphenols. Gorelik and associates 42 have shown that red wine may be equally important in collectively contributing to the polyphenols lower levels of postprandial cytotoxic lipid reduction of the risk of CVD. These factors appear to result from peroxidation products (malondialdehyde) in humans, another alcohol, from polyphenols present in some bever ages (especially possible mechanism for the benefits of the consumption of wine on wine), or from a combination of both, resulting in reduced risk of disease. thrombosis. 78

The early studies of Serge Renaud and his colleagues pointed out the importance of platelet function in the development of atherosclerosis, thrombosis, and CHD. 79 In an intervention study in humans, Zhang and coworkers 80 showed that alcohol, at physiologically relevant doses, has a dose dependent inhibitory effect on platelet aggregation. These authors concluded that their

findings are consistent with the view that alcohol reduces platelet sensitivity to thrombotic stimuli by inhibition of arachidonic acid release and there is subsequent thromboxane synthesis. Inhibition of platelet aggregation or function has also been shown by others. 81,82 Ruf 83 suggested that wine has a greater effect on platelet function than alcohol alone does, stating that the polyphenols in wine provide further reduction in prostanoid synthesis from arachidonate and also decrease platelet activity mediated by nitric oxide. Demrow and colleagues 84 demonstrated that polyphenols from both red wine and grape juice favorably affect platelet aggregation.

Many of the effects of alcohol and polyphenols on coagulation are transient effects, and the beneficial effects may last only 24 to 36 hours after someone has consumed alcohol. Furthermore, after heavy alcohol intake, there may be a rebound with increased platelet aggregation if further alcohol is not consumed, 83,85,86 although such rebound may be less after drinking wine than after consuming other beverages. 83 One explanation for the much lower CHD rates in France than in most other countries is that the French have traditionally consumed some alcohol (primarily red wine) with their evening meal every day. Thus, their clotting mechanisms Americans and northern Europeans tend to drink only on the weekend, often consuming a large number of drinks rapidly, a very

Additional mechanisms for protection of alcohol and polyphenols against CHD and myocardial infarction include decreased myocardial ischemia-reperfusion injury, 87,88 increased endothelial cell-dependent vasorelaxation, 89-92 simultaneous activation of endothelial cell antiapoptotic and proapoptotic pathways, 93 decreased plasma levels of factor VII 94 and fibrinogen, 95 increased fibrinolysis 96-102 and upregulation of fibrinolytic protein gene transcription in cultured human endothelial cells, 103,104 and increased levels of atrial natriuretic peptides. 105

There is also strong experimental evidence for additional mechanisms by which alcohol and wine polyphenols affect the initiation or progression of atherosclerotic lesions. These include The first recognized mechanism by which alcohol may prevent reduced LDL oxidation and aggregation, foam cell formation, and increased hsCRP of 0.32 (95% CI, 0.14-0.74) for consumers of 20 to in consumers of 5 to 7 drinks/week.

Booyse and coworkers 102 have demonstrated the sequence of molecular events by which endothelial cell expression of tissue plasminogen activator is increased through common activation of

of alcohol on HDL-C, fibrinogen, and hemoglobin A1c (the last a data lend support to a role of polymorphisms ADH1B and ALDH2 marker of glycemic control and insulin resistance) can almost combined with alcohol consumption in cancer, but other available entirely explain the reduced CHD risk among moderate drinkers. data are insufficient or inconclusive.

Hypertension

One cardiovascular risk factor that is generally not associated favorably with alcohol consumption is hypertension, and most alcohol and polyphenols may be cardioprotective (as well as studies show an elevation of blood pressure with increasing alcohol neuroprotective). In Figure 18-2, mechanisms of cardioprotection intake. 118,119 Some studies 120-122 show that the effect is mainly from by both alcohol and resveratrol, one of the polyphenols in red wine heavier drinking and that light alcohol intake has no effect or even that has been studied extensively, are illustrated. a slight lowering of blood pressure. For example, Klatsky and associates 121 found increases in blood pressure only among subjects colleagues 133 have carried out a number of intervention studies reporting three or more drinks per day, a finding similar to the - looking at the effects of wine on CVD risk factors. Their results association in all groups except African American men in an ARIC show that moderate wine consumption may result in an increase in report. ¹¹⁹ In the Luebeck Blood Pressure Study in Germany, Keil HDL-C, a decrease in the omega-6/ ¹ omega-3 ratio, and in some and associates 123 reported increases in blood pressure for men cases a slight increase in triglyc eride levels. Observed changes in consuming more than 40 g of alcohol per day (more than three hemostasis include reduced coagulation and increased fibrinolysis; typical drinks) or women reporting more than 20 g/day. Klatsky effects on blood pressure sure have been inconsistent. They have and associates 124 have shown that hypertension remains a strong found a reduction in inflammatory markers and an improvement risk factor for CVD regardless of alcohol intake, and hypertensives in endothelial function. These investigators concluded that who are heavy drinkers snow a substantial decrease their intake. 125 Whereas a direct typical US drinks) for men and nan unat dose for men association between alcohol and blood pressure has been necessary to prevent negative changes in cardiovascular risk association between alcohol and blood pressure has been necessary to prevent negative changes in cardiovascular risk association between alcohol and blood pressure has been necessary to prevent negative changes in cardiovascular risk association between alcohol and blood pressure has been necessary to prevent negative changes in cardiovascular risk association. who are heavy drinkers show a substantial decrease in blood limitation of consumption to below 30 g of alcohol daily (about 2 1/2 (notably CHD and ischemic stroke) show an inverse association hyperhomocysteinemia. Other clinical trials have demonstrated with moderate drinking.

Genetic Risk Factors

There is no question that individuals differ markedly in their response to alcohol consumption, for its beneficial and adverse health effects. Davey Smith 126 has described how "mendelian randomization – the random assortment of genes from parents to EFFECTS OF ALCOHOL ON OTHER TYPES OF offspring that occurs during gamete formation and conception – provides one method for assessing the causal nature of some environmental exposures." Such an approach avoids many of the The third leading cause of death in the United States, and the problems of confounding in observational studies. He concluded leading cause of disability, is stroke. It has been shown in many that "mendelian ran domization provides new opportunities to test studies that the risk of stroke related to ischemia (causality and demonstrates how investment in the human genome atherosclerosis), which is the type of stroke in about 80% of cases project may contribute to understanding and preventing the in the United States, is reduced by moderate drinking. 136,137 adverse effects on human health of modifiable exposures."

poorly understood. The most studied individual genes are those consuming five or more drinks per day. Others have shown that that relate to the metabolism of alcohol, especially those that control the risk of peripheral artery disease is reduced among moderate alcohol dehydrogenase (ADH) and acetal dehyde dehydrogenase drinkers. 135 (ALDH). Hines and coworkers 127 showed that moderate drinkers who are homozygous for the slow-oxidizing ADH3 allele had heart failure, recent scientific evidence suggests a lack of harmful higher HDL levels and a substantially decreased risk of myocardial effects from moderate alcohol consumption. Ettinger and infarction. 128 However, other studies have failed to show such an colleagues 140 described the "holiday heart syndrome" in 1978. association. 129 Jensen and coworkers 130 have found conflicting This syndrome often includes atrial fibrillation; the syndrome is results across different cohorts in their study relating lipoprotein usually not associated with long-standing heart disease and tends lipase to CHD.

the effects of alcohol on the risk of CHD; each tends to play only a syndrome, but others 144,145 have found no effect on the risk of small role by itself but a potentially large role in combination with atrial fibrillation for moderate alcohol intake, only for heavy other genes and environmental factors. Furthermore, certain genes drinking. A recent extensive review of experimental, clinical, and seem much more important in certain ethnic groups and cultures epidemiological data did not find evidence that alcohol is a factor, than in others. The large number of collaborative genome-wide certainly not a major factor, in the development of atrial association studies now being done, with huge numbers of subjects, fibrillation. 146 should help clarify the role that genetic factors play in the potential protection against cardiovascular and other diseases from alcohol use.

Similarly, for cancer, data on genetic factors modifying 305 alcohol's effects are unclear. Druesne-Pecollo and coworkers 131 published an overview of studies on the combined effects of alcohol drinking and polymorphisms in genes for ADH, ALDH, cytochrome P-450 2E1, and methylenetetrahydrofolate reductase

In some epidemiological studies, the combined beneficial effect on the risk of alcohol-related cancer. They concluded that current

Summary of Mechanisms

Collins and associates 132 have provided a summary of epidemiological and mechanistic studies demonstrating how

Clinical trials in humans are limited, but Leighton and improvements in glucose metabolism from the administration of alcohol. 67,134,135 In the future, more such trials as well as genetic studies will help determine the extent to which the beneficial effects of alcohol consumption on CVD risk seen in most observational studies are caused by the alcoholic beverage itself.

CARDIOVASCULAR DISEASES

Moreover, Klatsky and associates 138 found that an increase in risk At present, however, the association of genes with CHD is of hemorrhagic stroke from alcohol is seen only in subjects

For two other cardiovascular conditions, atrial fibrillation and to resolve when alcohol consumption is stopped. 141,142 Nissen and Current data suggest that there are many genes contributing to Lemberg 143 stated that even moderate drinking can lead to this

Reduces ischemia/reperfusion injury _

aggregation

♦ Prevents platelet

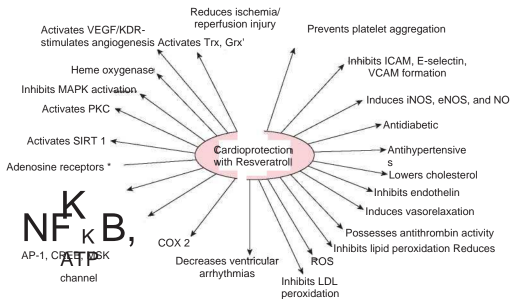


FIGURE 18-2 Cardioprotection from alcohol and from resveratrol. A, Cardioprotective and molecular targets of alcohol. Cardioprotective targets are shown on the right; the molecular targets are shown on the left. B, Cardioprotective and molecular targets of resveratrol. Cardioprotective targets are shown on the right; the molecular targets are shown on the left. (From Collins MA, Neafsey EJ, Mukamal KJ, et al: Alcohol in moderation, cardioprotection, and neuroprotection: epidemiological considerations and mechanistic studies. Alcohol Clin Exp Res 33:206, 2009. Reprinted with permission.)

cardiomyopathy, causing heart failure. It was previously believed the Survival And Ventricular Enlargement (SAVE) trial in subjects that heart failure, in general, would be made worse by any alcohol after myocardial infarction, Aguilar and coworkers 155 did not show consumption. As reviewed by Djousse and Gaziano, 147 however, a significant effect of moderate drinking on the development of most studies have shown that moderate drinkers are at lower risk heart failure or on survival. for developing heart failure. 148-153 For example, in a population of more than 100,000 subjects, Klatsky and associates 151 found that alcohol drinking was inversely related to risk of heart failure EFFECTS OF ALCOHOL ON DIABETES related to CHD (eg, at one or two drinks per day; RR, 0.6; 95% CI, AND METABOLIC SYNDROME 0.5-0.7), with consistency across subgroups of age, gender, ethnicity, education, smoking status, interval to diagnosis, and presence or absence of baseline heart disease or systemic hypertension. For heart failure not associated with coronary disease, moderate drinking was inversely related only in subjects who had diabetes mellitus. 151

For patients who already have depressed myocardial function, Cooper and associates 154 showed in the Studies of Left Ventricular Dysfunction (SOLVD) that moderate drinkers had lower

It is known that excessive alcohol intake can result in alcoholic subsequent all-cause mortality than abstainers did. In contrast, in

A major risk factor for the development of CVD not discussed before relates to glucose metabolism, especially the development of diabetes or the metabolic syndrome. Not only do data

support lower risk for development of these conditions, but for Effects of Alcohol Intake on Cardiovascular persons with diabetes or the metabolic syndrome, moderate drinking may lower their subsequent risk of CVD.

Development of Diabetes

Stampfer and colleagues 156 reported in 1988 that moderate drinkers 14 + g/day, in comparison with non drinkers. Solomon and in the Nurses' Health Study had a lower risk for developing colleagues 169 reported from the Nurses' Health Study that the risk diabetes than did abstainers; compared with nondrinkers, women for development of CHD among type 2 diabetics who were consuming 5 to 14.9 g of alcohol per day had an age-adjusted moderate drinkers was much lower than that for nondrinking relative risk of diabetes of 0.4 (95% CI, 0.3-0.6); for 15 g or more per diabetic subjects: for < 12 drinks daily, the relative risk was 0.72, day, the relative risk was 0.3 (95% CI, 0.2-0.4). The authors reported and for > 12 drinks daily, the relative risk was 0.45. In the that a strong inverse association between alcohol drinking and Physicians' Health Study, increasing levels of alcohol intake were body weight explained much of the apparent protective effect of associated with an even greater reduction in deaths from CHD alcohol. After simultaneous adjustment for Quetelet index (weight among diabetics than among nondiabetic subjects. 170 [kg]/height [m] ²), family history of diabetes, total calorie intake, and age, the relative risk of diabetes for consumers of 5 to 14.9 risk of CHD is 34% to 55% lower among diabetics who are g/day was 0.8 (95% CI, 0.6-1.2), and for women who drank 15 + moderate drinkers than in those who consume no alcohol. Among g/day, the relative risk was 0.6 (95% CI, 0.3-0.9). Reports in the more than 38,000 diabetics observed by the Kaiser ¹ Permanente British Medical Journal in 1995 from two large prospective studies group in California, those who consumed alcohol had evidence of 157,158 similarly showed a lower risk of diabetes among moderate considerably better control of their diabetes than did nondrinkers drinkers. Since then, many epidemiological studies 159-161 have 172; the authors concluded that this finding "supports current presented data supporting such an inverse association. Howard clinical guidelines for moderate levels of alcohol consumption and coworkers 162 have provided a good summary of the research among diabetes patients." on alcohol and diabetes, giving an estimate of 33% to 56% lower incidence of diabetes for consumers of one to three drinks per day. sequelae of diabetes but also the microvascular complications of A meta-analysis by Koppes and associates 163 indicated that for a diabetes. In a report from the EURODIAB Prospective wide range of alcohol intake (from about 12 to more than 3 Complications Study, involving the follow-up of 3250 type 1 drinks/day), the relative risk of diabetes for drinkers is about 30% diabetic patients from 16 different European countries, Beulens lower than it is for abstainers.

diabetes. Some studies have shown lower glucose levels and consumed alcohol moderately in comparison with nondrinkers. HbA1c. 164 In an intervention trial, Davies and coworkers 165 found The association was strongest among wine drinkers, and to some among nondiabetic postmenopausal women that the consumption extent among beer drinkers , but was not seen among those of 30 g/day of alcohol reduced fasting insulin concentration by consuming spirits. 19.2%, reduced triglyceride concentration by 10.3%, and increased insulin sensitivity by 7.2% but did not affect plasma glucose levels. epidemiological studies and limited clinical trials strongly Kroenke and associates 166 found that HbA1c was inversely suggests that moderate drinkers are less likely to develop associated with alcohol intake; among overweight women, there diabetes and its sequelae, especially CHD. Although not all of the was also an inverse association with insulin. Specifically, these - mechanisms are known, many studies suggest improved insulin investigators found that insulin levels were lowest for drinkers resistance or increased adiponectin levels. consuming no more than two drinks per day up to three days per week. In a large heterogeneous group of nondiabetic subjects, Development of Metabolic Syndrome insulin sensitivity was lower among abstainers than in all categories of drinkers. 167 Adiponectin serum levels were higher in The findings relating alcohol to metabolic syndrome show men consuming alcohol on 2 + days/week than in nondrinkers or associations with alcohol similar to those for diabetes. In the occasional drinkers; among women, those consuming all levels of $\,$ NHLBI Family Heart Study 174 and in NHANES III, 175 we found alcohol had higher adiponectin concentrations than nondrinkers evidence that moderate drinking is associated with lower risk of did. 167

intake to diabetic mechanisms. Sierksma and coworkers 134 carried association between estimates of lifetime alcohol intake and the out a randomized crossover study of 23 healthy middle-aged men metabolic syndrome. They found that subjects who averaged who were given 40 g of ethanol (four glasses of whiskey, versus no more than one drink per day for women or two drinks per day alcohol) on a daily basis for 17 days. They found that alcohol for men had an increase in their prevalence ratio for metabolic increased plasma adiponectin levels by 11% and increased the syndrome in comparison with drinkers who did not exceed these insulin sensitivity index in an insulin-resistant subgroup by 21%. A limits; nondrinkers were not included in these analyses. In a randomized crossover trial by Joosten and coworkers 67 showed meta-analysis on alcohol intake and metabolic syndrome, based that moderate alcohol consumption increased insulin sensitivity on data from seven previous studies with a total of 22,000 and ADIPOQ expression in postmenopausal women.

formerly abstaining diabetics who were randomly assigned to < 20 g of alcohol per day for women) was associated with 16% consume 5 ounces of either sauvignon blanc or merlot at dinner lower risk of metabolic syndrome for men and 25% lower risk for every evening or advised to continue to avoid alcohol. Significantly women; no significant effects were seen for heavier drinking. 177 lower levels of fasting blood glucose concentration during a 3month intervention period were seen among subjects consuming wine than were seen

for abstainers. The effect of the wine was variable, with **beneficial** effects primarily among those with more severe disease.

Disease Among Diabetics

Valmadrid and coworkers 168 reported that among persons with older-onset diabetes, the relative risk for death from CHD was 0.44 for subjects consuming 2 to 13 g/day and 0.21 for those consuming

A meta-analysis by Koppes and associates 171 indicates that the

Alcohol has been shown to affect not only the macrovascular and colleagues 173 found a lower occurrence of retinopathy, Mechanisms are unclear for the effects of alcohol intake on neuropathy, and nephropathy among those individuals who

In summary, evidence from a large number of prospective

most components of the metabolic syndrome (all components There have also been a number of clinical trials relating alcohol except for hypertension). Fan and coworkers 176 evaluated the subjects, Alkerwi and coworkers 177 found that the moderate Shai and colleagues 135 carried out a randomized trial among 91 intake of alcohol (defined as < 40 g of alcohol per day for men and

- Avoid obesity (keep the BMI <25).
- Consume a healthy diet, high in fiber and unsaturated fat and low in trans-fat and glycemic load (eg, a Mediterranean-type diet).
- Engage in moderate to vigorous physical activity (for at least 12 hours/day).
- Avoid smoking
- Unless contraindicated, consume a small amount of an alcoholic beverage regularly.

ALCOHOL AS A COMPONENT OF A HEALTHY 18 LIFESTYLE

Alcohol consumption, moderate or otherwise, should not be viewed in isolation but as part of broader social, cultural, and lifestyle issues. We now have good scientific data on which to base our definition of what constitutes a "healthy" diet and lifestyle. The Nurses' Health Study, the Health Professionals Study, and other research have defined a lifestyle that will lead to > 70% fewer myocardial infarctions and > 90% fewer cases of diabetes. This healthy lifestyle is shown in Box 18-2, based on research by Stampfer, ¹⁷⁸ Hu, ¹⁷⁹ Mukamal, ¹¹ and others.

Akesson and colleagues, 180 in a large prospective study among middle-aged and older women in Sweden, found that those subjects who met all five components of a healthy life style (defined as following what can be described as a Medi terranean-type diet, not smoking, not being obese, getting regular exercise, and consuming at least 5 g/day of alcohol) had a dramatically reduced risk of having a myocardial infarction. The authors suggested that if all women in their population had such a lifestyle, there would have been 77% fewer cases of CHD.

Khaw and associates, 181 in a large prospective observational study from the United Kingdom, assessed the effects on mortality of four "healthy behaviors": not smoking, being active, having evidence of a high fruit and vegetable intake, and consuming some alcohol but not more than 14 drinks per week (about 9X typical drinks by US standards). Increasing numbers of these behaviors were associated with lower risk of death from all causes as well as from CVD, cancer, and noncardiovascular causes. The authors calculated that the effect of these four healthy behaviors on total mortality, compared with none of them, is equivalent to a 14-year age difference in mortality risk.

Whereas some suggest that we should focus on the first four components of the healthy lifestyle and not be eager to encourage alcohol use, it has been shown that even among very healthy subjects (ie, who are lean, eat a healthy diet, are active, and do not Bone Mineral Density and Hip Fracture smoke), those who also consume a little alcohol have a much lower risk of heart disease and death. To address the issue of residual confounding by healthy lifestyle among drinkers, Mukamal and coworkers 182 restricted analysis to 8867 healthy men in the Health which has always assumed that elderly who drink alcohol may be Professionals Study who adhered to the first four low-risk behaviors (not including alcohol) and examined the association for sustaining fractures.) Mukamal alcohol consumption with CHD. In this group of "very healthy" men, there were still 106 incident CHD cases; the men who drank moderately (15.0 to 29.9 g/day) had a relative risk of 0.38 (95% CI, 0.16-0.89) compared with abstainers. This strong inverse association between moderate alcohol consumption and CHD in predominantly healthy individuals adds further evidence to support the hypothesis that the inverse association is causal and not confounded by healthy lifestyle behaviors. ¹⁰ On the other hand, a

paper with a restrictive definition of "healthy" reported no significant effects of alcohol on the risk of CHD among the healthiest subjects. 183

EFFECTS OF **ALCOHOL** ON NONCARDIOVASCULAR DISEASES

Before the encouragement of moderate drinking for the possible reduction in the risk of CVD is even considered, the potential effects on other diseases must be taken into account. In addition to CVD, it has long been known that moderate drinkers are at lower risk of gallstones. 184 There is interest in new data relating alcohol intake to obesity, bone mineral density, and dementia. As will be discussed, an increase in the risk of breast cancer from alcohol is of great concern for women, but recent data indicate a lower risk from moderate alcohol intake for certain other cancers. 185-187

Obesity

An unexpected finding in epidemiological studies is that obesity is often found to be less common among moderate drinkers than among abstainers. In the prospective Nurses' Health Study, 188 it was found that light to moderate drinking was not associated with weight gain in women (overall, a 16% lower risk for those reporting 15.0 to 29.9 g of alcohol per day versus abstainers). This potentially beneficial effect on weight gain was not seen in African American women or in heavier drinkers. Arif and Rohrer 189 evaluated alcohol intake and BMI among 8236 nonsmoking respondents who participated in the Third National Health and Nutrition Examination Survey. Current drinkers who reported drinking one drink or two drinks per day had 0.46 (95% CI, 0.34-0.62) and 0.59 (95% CI, 0.41-0.86) the odds of obesity, respectively, than did

Tolstrup and colleagues 190 found that among men, the odds ratios for having a high BMI among subjects drinking 1 to 3 days/month, 1 day/week, 2 to 4 days/week, 5 or 6 days/week, and 7 days/week were 1.39 (95% CI, 1.36-1.64), 1.17 (1.02-1.34), 1.00 (reference), 0.87 (0.77-0.98), and 0.73 (0.65 0.82), respectively. Similar associations were found for waist circumference, and corresponding results were found for women. These authors concluded that for a given level of total alcohol intake, obesity was inversely associated with drinking frequency, whereas the amount of alcohol intake was positively associated with obesity. Such an association was supported by Breslow and Smothers 191 among 45,896 never-smoking adults in the 1997-2001 National Health Inter view Surveys in the United States. As shown in Figure 18-3, from that study, more frequent drinking was associated with lower BMI. On the other hand, given drinking of a certain frequency, the number of drinks per drinking day was directly associated with

These results suggest that the frequent consumption of small amounts of alcohol is the optimal drinking pattern associated with a lower risk of obesity.

A number of epidemiological studies have shown that moderate drinkers have less osteoporosis and a lower risk of hip fractures than do abstainers. 192 (This goes against "conventional wisdom," more unsteady on their feet and at increased risk of falling and

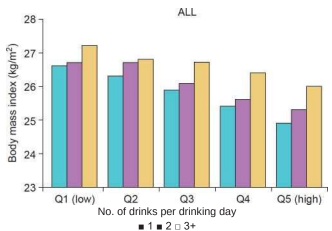


FIGURE 18-3 Alcohol consumption and BMI by frequency of drinking quintiles and quantity of alcohol. Association between alcohol consumption and BMI in stratified analyzes of frequency quintiles within quantity categories, National Health Interview Surveys, 1997-2001. Q, quintiles. (From Breslow RA, Smothers BA: Drinking patterns and body mass index in never smokers: National Health Interview Survey, 1997-2001. Am J Epidemiol 161:368, 2005. Reprinted with permission.)

and colleagues ¹⁹³ found that among older adults, moderate alcohol consumption had a U-shaped relationship with risk of hip fracture reported additional mechanisms for neuroprotection, as described but a graded positive relationship with bone mineral density at the earlier (under mechanisms of alcohol's effects on CVD). hip. A meta-analysis by Berg and coworkers 194 involving the review of 33 previous studies supports a reduced risk of hip fracture and higher bone density among men and women who consume small to moderate amounts of alcohol (in comparison with nondrinkers). There is probably a J-shaped curve, with ALCOHOL AND CANCER increased risk of fractures among heavier drinkers.

crazy

Among the most exciting recent scientific findings are that moderate drinkers tend to be less likely to develop Alzheimer's diseases, and these include alcoholic car diomyopathy, cirrhosis of disease and other types of dementia. This was reported for wine the liver, neurological diseases, and certain cancers. Of particular drinkers in Bordeaux by Orgogozo and colleagues in 1997, 195 and concern, however, are some conditions that may relate to even Truelsen and coworkers ¹⁹⁶ reported similar protection from wine moderate drinking, especially breast cancer in women. in Denmark, as did Panza and coworkers ¹⁹⁷ in a summary paper.

Britton and coworkers 198 reported from the Whitehall Study in the United Kingdom that there was less cognitive impairment in drinkers for tests of memory (borderline significant), verbal and abstaining men and women generally had a much greater risk of cognitive dysfunction than did occasional drinkers.

Espeland and colleagues 199 reported from the Women's Health Initiative Memory Study that compared with no intake, intake of one drink or more per day was associated with higher baseline Modified Mini-Mental State Examination scores (P < 0.001) and an odds ratio of 0.41 (95% CI, 0.23-0.74) for significant declines in cognitive function. Stampfer and coworkers 200 reported that among women who were moderate drinkers, compared with nondrinkers, the relative risk of impairment was 0.77 for general cognition (95% CI, 0.67-0.88) and 0.81 on the global cognitive score (95% CI, 0.70-0.93). Zhang and coworkers 201 reported that education modifies the putative effects of alcohol on memory, with better results among those with higher education, probably because of more moderate drinking patterns.

Ganguli and colleagues studied 202 men and women aged 74.4 years at baseline who were observed during 7 years with repeated assessments of cognitive functioning; abstainers showed lower baseline scores and a tendency for more rapid deterioration in cognitive functioning than did drinkers. Solfrizzi and colleagues 203 also found less progression from mild cognitive impairment to dementia for light drinkers in comparison with nondrinkers.

Mehlig and coworkers, ²⁰⁴ in a well-done analysis from a longterm prospective study of women in Sweden, reported that the lifetime risk of dementia was associated differently with wine consumption (a decrease in risk of dementia of 70%) and spirits consumption (an increase in dementia risk of about 50%). The authors concluded that the different associations by type of beverage suggest that the nonalcohol components in wine may be an important factor in lowering the risk of dementia. Whereas the investigators adjusted for the usual risk factors for dementia, including education and social class, there is always the possibility that other lifestyle factors that are different between wine drinkers and spirits drinkers may have influenced the results.

Peters and associates ²⁰⁵ carried out a meta-analysis on alcohol and dementia and concluded that moderate drinkers, especially wine drinkers, were at lower risk than abstainers for the development of Alzheimer's disease or other dementia. The mechanism for such protection is not known but may relate, among other factors, to prevention of cerebral atherosclerosis or decreased inflammation in brain tissue. Collins and associates 132 have

ADVERSE HEALTH EFFECTS **DRINKING:**

Most of the adverse health effects associated with alcohol are from drinking too much, too fast, or at an inappropriate time (as just before driving an automobile). Chronic heavy drinking can lead to a number of diseases that are often referred to as alcohol-related

Alcohol-Related Cancers

The strongest association between alcohol drinking and cancer is mathematical reasoning, verbal meaning, and verbal fluency. For for upper aerodigestive cancers, usually with an increased risk for men, the risk of dysfunction was lower by about 40% to 50% for a combination of heavy drinking and smoking. Weikert and drinkers, with the lowest risk at > 241 g/week of alcohol, the associates 206 reported that in comparison with drinkers averaging equivalent of about three drinks per day by US standards. Women no more than 6 g of alcohol (about 12 drinks) per day, squamous drinkers had lower point estimates for most measures, but upper aerodigestive cancers were increased among both men and statistically significant results were seen for verbal and women reporting > 30 g/day of alcohol (more than about 2 1/2 mathematical reasoning (OR = 0.3; CI, 0.2 -0.6) and for verbal drinks), with larger increases for heavier-drinking men. Allen and fluency (OR = 0.5; CI, 0.3 0.9), both at 49 to 80 g of alcohol per week coworkers, 207 in a study based on more than one million women in (the equivalent of about four to seven drinks per week). Lifetime the United Kingdom, found that for most cancers, the relative risk was lowest among women who reported up to two drinks per week (10 g of alcohol being considered a drink in this study) than among current abstainers, but this study was not

310 able to separate ex-drinkers from lifetime abstainers. Among drinkers, the strongest positive association between increased alcohol consumption and disease was for upper aerodigestive cancers (mouth, pharynx, esophagus), but this association was seen only among drinkers who were also smokers. In this study, lesser effects were noted for cancers of the rectum and breast.

Most studies show little effect of alcohol drinking on geni turinary tumors 185,208 and an apparent inverse association with renal cell carcinoma. 185,186 Furthermore, moderate drinkers seem to have lower risks of some types of leukemia ¹⁸⁷ and non-Hodgkin and other lymphomas. ^{209,210} In their large study, Allen and coworkers ²⁰⁷ showed among drinkers that an increase in alcohol intake of 10 g/day was associated with lower risks of thyroid cancer (RR = 0.75; 95% CI, 0.61-0.92), renal cell cancer (RR = 0.88; 95% CI, 0.78-0.99), and non Hodgkin lymphoma (RR = 0.87; 95% CI, 0.81-0.95). Other studies have shown that heavy drinking, but not moderate consumption, may increase the risk of pancreatic cancer. ²¹¹ A meta-analysis by Genkinger and colleagues 212 showed a slight positive association with pancreatic cancer risk for alcohol intake of 30 g/day (about 2 1/2 typical drinks) or more versus no alcohol (pooled multivariate relative risk, 1.22;

¹⁸ 95% CI, 1.03-1.45).

In an editorial on the association of alcohol intake with Barrett's esophagus and esophageal adenocarcinoma, El Serag and Lagergren ²¹³ described three population-based studies that demonstrated no overall effect of alcohol intake on either disease. Instead, these studies showed that the modest intake of wine, but not of other beverages, was associated with a significantly lower risk for both conditions.

Alcohol and Breast Cancer

Particular attention has been focused on the association between alcohol intake and the risk of breast cancer in women. The majority of epidemiological studies support the findings of Willett and coworkers 214 in 1987 and Longnecker ²¹⁵ in 1994: breast cancer risk is greater for women who consume alcohol than for abstainers. A notable exception is the Framingham Study, in which the long-term risk of breast cancer was not increased among drinkers in comparison with nondrinkers during a follow-up period of several decades. ^{216,217} An initial report from the Women's Health Initiative-Observational Study (WHI-OS) 218 describes a slight non dosedependent increase in risk of breast cancer for consumers of alcohol. In comparison with no alcohol, the adjusted relative risk for up to 5 g of alcohol per day (less than 12 drinks) was 1.10 (95% CI, 0.97-1.24); for 5 to 15 g/day, the relative risk was 1.14 (95% CI, 0.99-1.31); and for > 15 g/day, it was 1.13 (95% CI, 0.96-1.32).

In meta-analyses, the estimated increase in risk of breast cancer for the average consumption of one drink per day is usually between 6% and 15%. Stronger associations appear to be more common in hospital-based case-control studies than in cohort studies or community-based case-control studies, in studies published before 1990 than in studies published later, in studies with shorter follow-up periods, and in studies conducted outside of the United States than in US studies. ²¹⁹

Some studies show that the increase in risk of breast cancer among women who drink alcohol may be prevented or ameliorated if the women are consuming adequate amounts of folate in the diet, ^{220,221} but this has not been supported by certain other studies. ^{218,222} In the WHI-OS, there was no effect of folate intake on the association between alcohol and breast cancer. ²¹⁸ Other epidemiological research suggests an increase in breast cancer risk from alcohol intake only among women who are also taking hormone replacement therapy ^{223,224} or those who binge drink. ²²⁵

Small amounts of alcohol are associated with a larger effect on the risk of CVD and other diseases of aging that are much more common causes of death; CHD deaths are nine times more frequent than breast cancer deaths, and stroke is three or four times more common. Hence, even for a slight increase in the risk of breast cancer for alcohol intake, on average a postmenopausal woman who stops any alcohol intake in an attempt to lower her risk of breast cancer may be increasing her risk of other diseases, and the net effect may be a shorter life span. In any case, advice about alcohol consumption for a postmenopausal woman with strong cardio vascular risk factors will be different from that for a young woman with no cardiovascular risk factors but a very strong family history of breast cancer.

ALCOHOL AND TOTAL MORTALITY

The bottom line is total mortality. Certain religious groups that limit the intake of meat and prohibit the use of alcohol and tobacco tend to have lower mortality rates. For the general population, alcoholics and binge drinkers do not live as long as abstainers, but regular, moderate drinkers live longer than abstainers. In a study of more than 270,000 men by the American Cancer Society, ²²⁶ the risk of dying of any cause was 16% lower for those reporting one drink per day than for abstainers. Thun and colleagues ²²⁷ evaluated alcohol and mortality among 490,000 Americans; all-cause mortality rates were 21% lower among men and women reporting about one drink daily than among nondrinkers. Doll ²²⁸ reported that British physicians who were moderate drinkers had lower total mortality rates than lifetime abstainers.

Di Castelnuovo and colleagues, ²⁴ in a meta-analysis based on more than one million subjects from 56 independent prospective studies, found that the relationship between alcohol intake and total mortality is J-shaped, with about 16% reduced risk for light drinkers and increased mortality for heavy drinkers, as shown in Figure 18-4. In the meta-analysis, differences were noted between men and women for the association between alcohol intake and mortality (Fig. 18-5).

As shown in Figure 18-5, the mortality risk for women drinkers exceeded that of nondrinkers at just more than 2 drinks per day, whereas the increase above that of abstainers for men was at about 3 122 drinks per day. 24 In subanalyses, the relative risk of drinkers exceeded that of nondrinkers at about 30 g of alcohol per day (the equivalent of about 2% drinks) in studies in the United States but only at about 65 g/day (or about 5 typical drinks) in studies in Europe. Whereas the reasons for this are not known, they may relate to differences in drinking practices, in that consuming alcohol on a regular basis and with meals is a usual pattern in much of Europe. Among women, the differences between studies in the United States and Europe were less marked. 24

Contribution of Moderate Drinkers to Alcohol-Related Total Mortality

Some studies claim that moderate drinkers make a large contribution to deaths attributed to alcohol. In a summary of the association of alcohol consumption with mortality in Canada for 2002, Rehm and coworkers ¹⁴ showed that when moderate drinking was based on an average weekly consumption of 14 or fewer drinks for men or 7 or fewer drinks for women, the number of deaths "caused" by alcohol exceeded those "pre vented" by alcohol, as shown in Figure 18-6A.

However, when subjects classified as moderate drinkers who reported episodes of binge drinking were excluded, there was a marked difference in the number of deaths attributed to alcohol, as shown in Figure 18-6B. Removal of binge drinkers from the moderate category lowered especially the

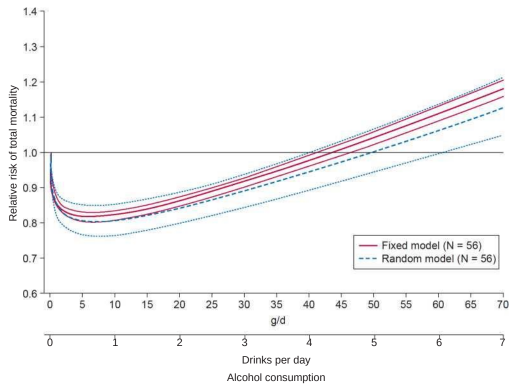


FIGURE 18-4 Alcohol intake and total mortality: a meta-analysis. Relative risk of total mortality (95% CIs) by alcohol intake, extracted from 56 curves using fixed- and random-effects models in a meta-analysis. (From Di Castelnuovo A, Costanzo S, Bagnardi V, et al: Alcohol dosing and total mortality in men and women: an updated meta-analysis of 34 prospective studies. Arch Intern Med 166:2437, 2006. Reprinted with permission.)

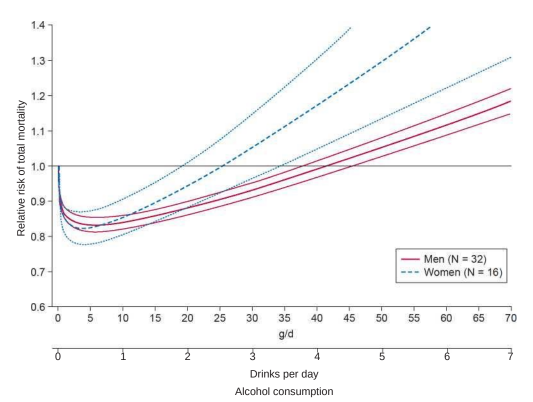
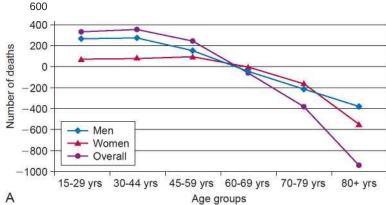
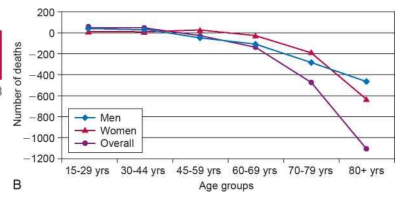


FIGURE 18-5 Alcohol intake and total mortality, by gender: a meta-analysis. Relative risk of total mortality (99% CIs) and alcohol intake in men and women in the United States, Europe, and other countries (Australia, Japan, and China), extracted from adjusted curves in a meta-analysis. (From Di Castelnuovo A, Costanzo S, Bagnardi V et al: Alcohol dosing and total mortality in men and women: an updated meta-analysis of 34 prospective studies. Arch Intern Med 166:2437, 2006. Reprinted with permission.)









number of excess deaths in younger people. Rehm and coworkers ¹⁴ concluded: "If moderate consumption is based on average volume alone, 866 net deaths in 2002 among those younger than 70 years of age were due to moderate consumption of alcohol (1.3% of all the deaths in this age group, consisting of 1653 deaths caused and 787 deaths prevented). When heavy drinking episodes were excluded, the net effect was beneficial (55 prevented deaths, 0.09% of all deaths); the net burden was higher for younger ages and the net benefits for older ages."

Differential Effects of Alcohol on Total Mortality According to Age

The diseases that tend to be "protected against" by moderate drinking occur later in life, among middle-aged and older adults. ¹⁴ There are few health benefits of drinking among the young (who, in any case, drink for the social effects of alcohol). Binge drinking, especially among the young, continues to be a serious public health problem around the world. In Germany, deaths caused by alcohol occurred mainly in young people (eg, from accidents) or middle-aged adults (eg, from cirrhosis), whereas most of the benefits on mortality were seen among the elderly. ²²⁹

There are an increasing number of reports describing the associations of alcohol use over the life span with health and disease. Powers and Young ²⁰ showed that middle-aged women in Australia who were moderate drinkers (who con sumed the equivalent of up to about 10 typical US drinks per week) had the best self-reported state of overall health. Furthermore, those who remained moderate drinkers through out the study continued to have the highest rating of health and self-reported higher quality of life; those who stopped drinking during follow-up showed a decrease in their ratings of overall health.

FIGURE 18-6 Number of net deaths attributable to moderate alcohol consumption. A, Number of net deaths in Canada in 2002 attributable to persons who reported alcohol intake, on average, placing them in the moderate drinking category, without consideration of binge drinking. Moderate drinking is defined as an average of < 40 g/day for men and < 20 g/day for women. B, Number of net deaths attributable to moderate drinkers who did not report any occasions of binge drinking (such drinkers were excluded from the analysis): sensitivity analysis. (From Rehm J, Patra J, Taylor B: Harms, benefits, and net effects on mortality of moderate drinking of alcohol among adults in Canada in 2000, Ann Epidemiol 17:S81, 2007. Reprinted with permission.)

ARE THERE DIFFERENCES IN EFFECT BY TYPE OF ALCOHOLIC BEVERAGE?

Most studies show beneficial effects of all types of alcoholic beverages—beer, wine, and spirits—on the risk of CHD and other diseases, but many show more favorable associations among wine drinkers. ²³⁰⁻²³³ Some studies have found wine drinkers to have a healthier diet overall than drinkers of beer or spirits, which may explain some of the putative additional beneficial effects of wine on health outcomes. ²³⁴⁻²³⁷ Still, even with adequate adjustment for diet and other lifestyle factors, many welldone observational studies (especially those among European populations) show significantly lower morbidity and mortality for wine drinkers than for consumers of other beverages. In the prospective ARIC study in the United States, King and associates ¹⁹ found that subjects initially reporting no alcohol intake who

reported moderate drinking at a later examination had a 38% lower risk for a cardiac event during the following 4 years; new wine drinkers had better health outcomes than did new drinkers of other beverages.

Streppel and colleagues ²³⁸ have reported that in a long -term follow-up study among men in the Netherlands (until death in most of the subjects), with repeated assessments of alcohol intake, moderate drinkers had lower rates of cardio vascular and all-cause mortality than did abstainers; fatality risks were lower for wine consumers than for consumers of other beverages. Analyzes of survival after the age of 50 years from that study, by type of beverage, are shown in Figure 18-7.

As shown in Figure 18-7, the consumption of any type of alcohol (versus not drinking) was associated with some what greater survival after the age of 50 years. The median survival after the age of 50 years was several years greater for wine consumers than for nondrinkers and greater than for consumers of other beverages. ²³⁸

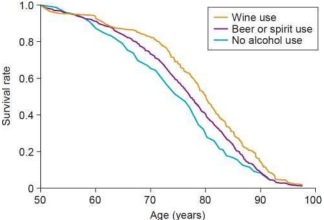


FIGURE 18-7 Survival after the age of 50 years according to long-term alcohol consumption. Survival curves for men with long-term consumption of alcohol from wine, beer, or spirits and for no alcohol consumers within the Zutphen Study. Adjusted for baseline energy intake without energy from alcohol; the number of cigarettes smoked; cigar or smoking pipe; intake of vegetables, fruit, fish, and saturated and trans fatty acids; body mass index; prevalence of myocardial infarction, stroke, cancer, and diabetes mellitus; and socioeconomic status. (From Streppel MT, Ocke MC, Boshuizen HC, et al: Long-term wine consumption is related to cardiovascular mortality and life expectancy independently of moderate alcohol intake: the Zutphen Study. J Epidemiol Community Health 63:534, 2009. Reprinted with permission.)

In a large study in Finland among older men of a similar high socioeconomic class (business executives) who were observed during 29 years, those who stated that they preferred wine (rather than beer or spirits) had lower, fully adjusted mortality rates (especially for CVD) and a higher quality of life among survivors compared with subjects reporting the intake of other beverages. ⁴⁹ These findings from epidemiological studies are strongly supported by a vast amount of experimental evidence showing that many of the polyphenols in wine have beneficial effects on biological and genetic mechanisms associated with the development of CHD and other diseases.

CONCLUSION

The scientific data from epidemiological studies, basic science, and limited clinical trials support a role for moderate alcohol consumption in the prevention of CVD. At the same time, giving advice about alcohol intake must take into consideration that this substance can be a "double-edged sword." ²³⁹ We have long known that excesses (in habits, foods, or alcohol) have problems that are inherent not in the activities or substances themselves but in their inappropriate use. For example, in an address to a temperance society in 1842, Abraham Lincoln stated: "It has long been recognized that the problems with alcohol in this country relate not to the use of a bad thing, but to the abuse of a good thing." ²⁴⁰ There are no data showing that encouragement of moderate consumption increases abuse, but it is clear that advice about alcohol should vary according to the characteristics of the individual patient.

As described by Cole, ²⁴¹ the finest moral rationale for prevention-oriented public health activity should be informing people, and it should not be based on "paternalism" ("we know what is best and will tell you only what you need to know"). We have had examples of sound scientific information relating alcohol intake to CHD that has been sacrificed to be "politically correct." ²⁴²

There are certain people who should not drink at all 313 (including former abusers of drugs or alcohol, people with

certain medical conditions, children and adolescents, and people with religious or moral proscriptions against alcohol), and there can never be a general recommendation for every body to consume alcohol. On the other hand, we should not withhold from our patients and the public scientifically sound and balanced data on alcohol and health. Whereas our current understanding suggests that moderate, sensible drinking can be potentially helpful for prevention of CHD in most adults (those without contraindications) as one component of a healthy lifestyle, any recommendations for its use in the individual should be based on consultation with the health care provider. Given the potential for misuse or abuse, however, major organizations such as the American Heart Association and the American College of Cardiology have not promoted or endorsed general recommendations aimed at the public at large.

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KEY POINTS

- Obesity is the leading public health crisis of our time. It is a primary target to reduce an avoidable disease burden in the United States.
- Obesity is an independent risk factor for major cardiovascular events, including coronary heart disease, heart failure, and stroke.
- Body mass index is a vital sign for assessment of patients with excess body weight and for stratification of treatments according to the likelihood of underlying disease risk.
- A series of metabolic abnormalities in obesity culminate in a clustering of risk factors for cardiovascular disease and type 2 diabetes mellitus, a condition known as the metabolic syndrome.
- Framingham risk score assessment for cardiovascular disease does not account for obesity. Risk assessment should include body mass index, waist circumference, blood serum biomarkers, and physical fitness.
- Major studies show that lifestyle change leading to weight loss can reduce or reverse risks associated with cardiovascular disease, including sleep apnea, hypertension, type 2 diabetes, and coronary heart disease.
- In patients intractable to lifestyle intervention alone, pharmacotherapy can facilitate weight loss to reduce risk of comorbid conditions. In cases of severe obesity, bariatric surgery is associated with significant weight loss and improvement in diabetes, hypertension, and obstructive sleep apnea.

CHAPTER 19

Overweight, Obesity, and Cardiovascular Risk

George L. Blackburn, Kristina Spellman, and Samuel Wollner

Obesity prevention and cardiovascular risk reduction will require a new approach that takes into account the sociopolitical,

Obesity is the leading public health crisis of our time. The most recent data from the National Health and Nutrition Examination Survey (NHANES 2005-2006) indicate that the prevalence of obesity (BMI > 30 kg/m¹² 13) among adults is 34% 1; for extreme obesity, the 2 6%. Minorities figure is disproportionately affected. Approximately 53% of non-Hispanic African American women and 51% of Mexican American women 40 to 59 years of age are obese compared with an estimated 39% of non-Hispanic white women of the same age. 2 Together, overweight and obesity affect more than 66% of the adult population.¹⁴

Rates of childhood obesity reflect those for adults. The proportion of 18- to 29-year-olds who were obese in 2004-2006 more than tripled from 8% in 1971-1974 to 24%. ¹ NHANES data for the combined years of 2003-2006 indicate that 16.3% of children and adolescents 2 to 19 years of age are obese, defined as at or above the 95th percentile of the 2000 BMI-for-age growth charts; 11.3% of children and adolescents in the same age group are above the 97th percentile, or

Many potential mechanisms have been proposed to explain the association of obesity with cardiovascular events, including increased severity of CVD, systemic inflammation, insulin resistance, neurohormonal activation, and abnormalities in adipokine pathways (Fig. 19-1). ¹⁰ Excess weight exacerbates a number of cardiovascular and metabolic risk factors (Box 19-1). ¹² Inflammatory adipokines may increase insulin resistance and diabetes, ¹³ and abnormal lipid metabolism, which is common in those with obesity, can lead to atherosclerosis- ¹⁴ rotic plaque. ¹⁴

Excess adipose tissue, especially intra-abdominal, 12 predisposes patients to type 13diabetes, hypertension, dyslipidemia, etc the metabolic syndrome largely through

extremely obese.15

Low-income and minority children, like adults, are disproportionately affected. ¹⁶ ¹⁷ The rates of obesity in Hispanic and non -Hispanic black children are 18.5% and 11.8%, respectively, compared with 12.6% in non-Hispanic white children. Evidence suggests that obesity-associated morbidity may increase with longer duration of the disease, ^{1,6} adding even more urgency to the need to reverse the trend and to reduce future morbidity. ¹

Modern therapies, such as statins, and lifesaving procedures have reduced the rate of mortality due to heart disease in the United States. There are, however, concerning trends in both non-fatal events and the disability that results from major cardiovascular events. ¹⁸Risk factor levels continue to rise in the United States at alarming rates. ¹⁹

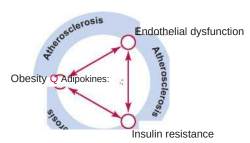
economic, and environmental forces that interact to create an obesogenic environment.

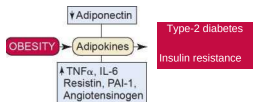
Addressing these risk factors will be the first step in alleviating avoidable burdens on patients and our health care system.²⁰

Obesity is an independent risk factor for cardiovascular events in the general population , patients with established cardiovascular disease (CVD), and elderly persons. ²¹ During the last decade, more than 100 prospective cohort studies and three meta analyzes (with more than 90 prospective studies and 1.1 million participants) ²²have confirmed obesity's central role in the development of CVD.

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increased lipolysis that raises the product tion of free fatty acids and adipokines. The excess fatty acids interfere with insulin receptor signaling and lead to decreased glucose transport, often referred to as lipotoxicity. They also activate protein kinase C (through increased fatty acyl coenzyme A and diacylglycerols). ¹³





Dyslipidemia

Endothelial dysfunction Hypertension

I AGE

Atherosclerosis

I Oxidative stress

FIGURE 19-1 Adipokines serve as the cellular mediators of metabolic syndrome and endothelial dysfunction. (From Lau DC, Dhillon B, Yan H, et al: Adipokines: molecular links between obesity and atherosclerosis. Am J Physiol Heart Circ Physiol 288:H2031, 2005.)

BOX 19-1 Risk Factors for Atherosclerosis and Vascular Disease Associated with Obesity

Hypertension

19

- Dyslipidemia
- Diabetes mellitus (type 2)
- Obstructive sleep apnea
- Hyperinsulinemia, insulin resistance
- Low levels of plasminogen activator inhibitor
- High levels of C-reactive protein
- Hyperviscosity
- Framingham risk score

signal transduction. Excess free fatty acids also impair phosphoinositide 3-kinase activation in response to insulin, leading to decreased activity of glucose transporter 4 (GLUT4), an important insulin-sensitive glucose transporter in muscle and fat. ¹³ Growing evidence suggests that insulin resistance in liver, muscle, and adipose tissue is associated with and may be the result of increased proinflammatory cytokines. ^{15,16}

In an obese state, some adipokines—proteins, such as tumor necrosis factor—a , or cytokines, such as interleukin-6—are elevated. Adipokines inhibit insulin action and contribute to proinflammatory effects, insulin resistance, and endothelial dysfunction. Adiponectin and resistin have recently been associated with incident heart failure. ¹⁷

This series of metabolic abnormalities culminates in a clustering of risk factors for CVD and type 2 diabetes mellitus, a condition known as the metabolic syndrome or prediabetes. The risk factors include raised blood pressure, dyslipidemia (increased triglycerides and lowered high-density lipoprotein cholesterol [HDL-C]), high fasting glucose concentration, and central obesity (Table 19-1). ¹⁸ Whereas obesity is an independent risk factor for CVD, ¹⁹ the complex interaction between excess body weight, lipid oxidation, and hyperglycemia underlies a strong connection between insulin resistance and risk for CVD. Efforts to prevent CVD must focus strongly on prevention of excess weight gain, insulin resistance, and dyslipidemia. ²⁰

Abdominal obesity	Waist circumference* Population- and country-specific definitions
	> 150 mg/dL
Triglycerides (drug treatment for elevated triglycerides is an alternate indicator ·)	
HDL-C (drug treatment for reduced HDL-C is an alternate indicator ·)	Men < 40 mg/dL Women < 50 mg/dL
	> 130/ > 85 mmHg
Blood pressure (hypertensive drug treatment in a patient with a history of hypertension is an alternate indicator)	
Fasting glucose - (drug treatment of elevated glucose is an alternate indicator)	> 100 mg/dL

- *It is recommended that the International Diabetes Federation cut points be used for non-Europeans and either the International Diabetes Federation or American Heart Association/National Heart, Lung, and Blood Institute cut points be used for people of European origin until more data are available.
- ^The most commonly used drugs for elevated triglycerides and reduced high-density lipoprotein cholesterol (HDL-C) are fibrates and nicotinic acid. A patient taking one of these drugs can be presumed to have high triglycerides and low HDL-C. High-dose omega-3 fatty acids assume high triglycerides.
- · Most patients with type 2 diabetes mellitus will have the metabolic syndrome by the proposed criteria.

OBESITY ASSESSMENT AND RISK OF CARDIOVASCULAR DISEASE

To stratify people in different demographic and ethnic groups according to their 10-year risk for coronary heart disease (CHD) events, clinicians use the Framingham risk score. ²¹ Guidelines recommend use of the Framingham risk score, or a modified version of it, to identify high-risk individuals (10-year risk > 20%) who can benefit from aggressive risk reduction measures. ²²⁻²⁴ Although the Framingham risk score is widely used and highly valuable, it does not include

TABLE 19—1 Criteria for Clinical Diagnosis of the Metabolic Syndrome

You can calculate BMI as follows

weight (kg) BMI = height squared (m²)

If pounds and inches are used weight (pounds) X 703

height squared (inches 2)

FIGURE 19-2 Diagnosis of overweight and obese: using body mass index (BMI). (From The Practical Guide: identification, evaluation, and treatment of overweight and obesity in adults , Bethesda, Md, 2000, National Heart, Lung, and Blood Institute, Department of Health and Human Services. NIH publication 00-

BOX 19-2	Standard Criteria for Body Mass Index
Underweight Healthy weight Overweight Obese Morbid obesity	BMI, kg/ ^{m2} 18.5 18.5-24.9 25.0-29.9 30-39.9 > 40

measures of obesity or inflammation. This limits its clinical relevance in assessing intermediate risk for CHD.

Body mass index (BMI) is an important screening tool to assess patients with excess body weight and to stratify treatments according to the likelihood of underlying disease risk (Fig. 19-2). ²⁵ BMI is calculated as weight in kilograms divided by height in meters squared (kg/m²) and categorizes obesity into three classes: class I, BMI of 30 to 34.9; class II, BMI of 35 to 39.9; and class III, BMI > 40, or extreme obesity (Box 19-2). Across genders and ethnicities, increased BMI is associated with greater comorbidity burden (Fig. 19-3). ²⁶ BMI may provide a better determination of global disease risk than weight alone, but it is of limited diagnostic value in very muscular individuals and those with little muscle mass, such as elderly patients. ^{25,27}

Waist circumference as a measure of abdominal or central obesity has attracted particular attention because of its inclusion as a prerequisite for the diagnosis of metabolic syndrome. 11,28 It provides important additional prognostic information, especially when an unhealthy level of excessive adiposity is suspected. 25,27 A higher risk for diabetes, dyslipidemia, hypertension, and CVD has been associated with a waist circumference > 102 cm (> 40 inches) in men and > 88 cm (35 inches) in women (although the International Diabetes Federation ¹⁸ has specified lower cut points of > 94 cm in men and > 80 cm in women for European whites and > 90 cm in men and > 80 cm in women for certain Asian populations and for those of central or South American ancestry; Table 19-2). A study showed that either BMI or waist circumference independently predicted or was associated with type 2 diabetes. 29

Unlike in definitions of obesity in adults, the growth curve needs to be taken into account in children. 13 The 2000 growth chart developed by the Centers for Disease Control and Prevention (CDC) is based on national height and weight data for children 2 to 19 years of age. The CDC and the National Heart, Lung, and Blood Institute (NHLBI) have defined obesity in children as a BMI > 95% on the 2000 growth chart and overweight as > 85%. These cutoff points are arbitrary and, unlike adult definitions of overweight and obesity, are not based on health risk data. 13

In the United States, 23 million adults with no history of CVD are classified as intermediate risk by the Framingham score (a 10year risk for major CHD events of 10% to 20%). 30 New or emerging risk factors, particularly inflammatory markers and markers of atherosclerotic burden, offer promise as screening tools for these individuals. ²⁴ However, current data from the US Preventive Services Task Force concludes that there is sufficient evidence to recommend the use of high-sensitivity C-reactive protein (hsCRP) among initially intermediate-risk persons. 31 At the discretion of the physician , hsCRP measurements may help direct further evaluation and therapy in primary prevention of CVD. 32

Childhood Obesity

Childhood obesity has wide-ranging comorbidities with clinical, psychosocial, and economic ramifications. 33 Children who are obese in their preschool years are more likely to be obese in adolescence and adulthood. 34 They are also more likely to develop diabetes, hypertension, hyperlipidemia, asthma, and sleep apnea. 34 Other risk factors for the development of CVD and type 2 diabetes in children include a sedentary lifestyle, family history of type 2 diabetes, high low-density lipoprotein cholesterol (LDL-C) levels, hyperinsulinemia, insulin resistance, 19 and high total cholesterol concentration. 35

With the worldwide epidemic of childhood obesity, disorders once mainly found in adults, such as metabolic syndrome, are occurring in children. ³⁶ The term *pediatric metabolic syndrome* includes a cluster of cardiovascular risk factors, such as insulin resistance, dyslipidemia (including increased triglycerides and decreased HDL-C), hypertension, and obesity. Children with metabolic syndrome have significantly higher BMI and glucose and triglyceride levels and lower HDL-C than do those without the syndrome. ³⁵ Prevalence of the metabolic syndrome in obese children is reported at 30%. 37

A study of 214 overweight and obese Costa Rican children up to 10 years of age found that obese children had lower mean serum levels of HDL-C and significantly higher mean serum concentrations of insulin, hsCRP, and triglycerides than their overweight peers did. They also had higher insulin resistance. 35

Maffeis and colleagues 38 tested 1044 Italian children aged 6 to 11 years for blood pressure, serum triacylglycerides, total glucose, cholesterol, HDL-C, insulin, and aminotransferase . The prevalence of high blood pressure in overweight boys and girls was 14.3% and 6.4%, respectively; in obese boys and girls, it was 40.4% and 32.8%, respectively. High blood pressure increased progressively with BMI z -score cat egories and waist-to-height ratio. Hypertensive children had significantly higher insulin and insulin resistance.

Adult Obesity

Because of its maladaptive effects on various cardiovascular risk factors and its adverse effects on cardiovascular structure and function, obesity has a major impact on cardiovascular diseases, such as heart failure, CHD, sudden cardiac death, and atrial fibrillation. ^{26,39} A large body of literature shows that obesity can cause and exacerbate many chronic diseases, such as diabetes, hypertension, dyslipidemia, stroke, and obstructive sleep apnea. ⁴⁰ It more than doubles the risk of heart failure. ⁴¹

Epidemiological data suggest a linear relationship between BMI and CHD. Jee and coworkers 42 conducted an analysis of



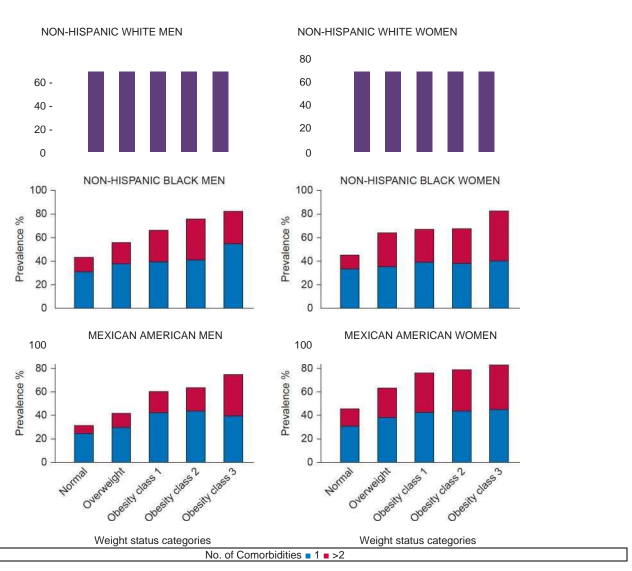


FIGURE 19-3 Prevalence of one and two or more overweight- and obesity-related morbidities by weight status category for sex, race, and ethnic subgroups. (From Must A, Spadano J, Coakley EH, et al: The disease burden associated with overweight and obesity. JAMA 282:1523, 1999.)

BMI and CHD incidence among 133,740 participants during 9 years of follow-up. The authors found that after adjustment for age, gender, and smoking status, each unit increase in BMI was associated with a 14% higher risk of incident CHD. Even a normal BMI of 24 to 24.9 was associated with a twofold increased risk of CHD.

A 20-year follow-up analysis of the Nurses' Health Study cohort showed a graded relationship between increasing BMI and incidence of CHD. ⁴³ Compared with normal-weight women, the relative risk of CHD in overweight women was 1.43. For obese women, it was 2.44. In the Women's Health Initiative, overweight and obesity were significantly associated with CHD incidence in both white and black women. ⁴⁴

The Framingham Heart Study, which observed more than 5000 individuals for up to 44 years, also reported substantial cardiovascular risk linked to overweight and obesity. ⁴⁵ One analysis found that overweight and obesity were independently associated with an increased risk for CVD as well as established risk factors, including hypertension, hypercholesterolemia, and type 2 diabetes. ⁴⁵ A prospective study of more than 17,000 healthy female US nurses with an average age of 50 years found that women who were obese at midlife were 79% less likely to be healthy at the age of 70 years compared with those who were lean in their 40s and 50s. The odds of being healthy among those who were obese at the age of 18 years and then gained more than

22 pounds by middle age were reduced by 82%. 46

RISK FACTORS

Visceral Adiposity

Data show that central obesity poses a more significant CVD risk than total obesity and that waist circumference ¹² and waist-to-hip ratio, common surrogates for abdominal or central obesity, may be better predictors of atherosclerosis and CVD risk than BMI. ^{39,47-51} Adipose tissue, especially intra abdominal visceral fat, has an independent endocrine function that leads to the release of inflammatory adipokines, including tumor necrosis factor- a , interleukin-6, and plasminogen activator inhibitor type 1. ^{12,52}

Inflammatory adipokines may increase insulin resistance and diabetes in obesity ⁵² and heighten risk for thrombosis. ¹² They may also affect the progression of endothelial dysfunction, further increasing inflammation and the risk for

birth weight and later development of visceral or central adiposity 55,57 as well as metabolic syndrome 55 and higher blood pressure later in life. 54,58-60 Blood pressure is influenced by size at birth 61,62 as well as by weight gain in childhood. 63 Abnormalities are accompanied by functional changes in the vascular tree, and evidence of early alterations in vascular function has been described in children and adolescents with low birth weight. 61.64

Hypertension

In 2003-2006, 36% of men and women between the ages of 45 and 54 years had hypertension compared with 65% of men and 80% of women 75 years of age and older. ¹ Large differences in blood pressure by ethnic group exist among adults. 65 The National Health Interview Survey found that in 2007, 23% of US adults had been told by a physician or health professional on two or more visits that they had hypertension. ⁶⁶ The condition in adults is associated with increased risk of myocardial infarction, stroke, and cardiovascular mortality. 67 Hypertension often increases with rising body weight. ^{68,69} It also affects 1% to 5% of children and adolescents. 70 The condition increases progressively with higher BMI and can be detected in approximately 30% of overweight children (BMI > 95th percentile). 71

In a cross-sectional study of 710 subjects aged 20 to 25 years, Dimkpa and Oij 69 found a significant correlation between BMI and systolic and diastolic blood pressure and resting heart rate after controlling for age and physical activity status. Overweight and obese subjects had a significantly higher risk of hypertension than non-overweight or obese controls did, and the prevalence of hypertension and tachycardia rose with increases in BMI. 69

Like adults, children and adolescents with severe elevation of blood pressure are at risk of adverse outcomes including cerebrovascular accidents and congestive heart failure. 67 Two autopsy studies in adolescents and young adults found significant relationships between the level of blood pressure and the presence of atherosclerotic lesions in the aorta and coronary arteries. 72,73 Childhood levels of blood pressure are also associated with carotid intima-media thickness, 74 large artery compliance, 75 decreased brachial artery flow-mediated vasodilation, 76 and left ventricular hypertrophy at a level associated with a fourfold greater risk of adverse cardiovascular outcomes in adults. 67 Overweight and high blood pressure are also components of metabolic syndrome, a condition of multiple metabolic risk factors for CVD as well as type 2 diabetes. 11 These outcomes underscore the need to prevent obesity early in life to protect against future life-threatening 69 consequences. 69

Sleep Apnea

Obstructive sleep apnea is independently associated with increased cardiovascular risk. 77 It has also been linked to insulin resistance and glucose intolerance and is independently associated with impaired glycemic control 77 and type 2 diabetes in patients who report excessive sleepiness. ⁷⁸ Obstructive sleep apnea is strongly correlated with intra- abdominal fat, 79,80 and serum lipid levels are elevated in patients who have obstructive sleep apnea. 81 Prospective findings from up to 15 years of followup data from the Wisconsin Sleep Cohort Study indicate that untreated sleep apnea predicts increases in blood pressure, hypertension, stroke, depression, and mortality. 82

Obesity causes and exacerbates obstructive sleep apnea, 40,80 and increases in weight have been associated with a rising prevalence of obstructive sleep apnea. Data show a fourfold rise in obstructive sleep apnea with each increase in the

	100				
	Current Recommende Abdominal Obesity by		nce Thresholds for		
Recommended Waist Circumferenc Threshold for Abdominal Obesity Men Women					
	Organization				
Europidus	IDF	> 94cm	> 80cm		
Caucasian	WHO	> 94cm (increased)	> 80 cm (increased risk)		
		> 102 cm (still higher risk)	> 88 cm (still higher risk)		
United States	AHA/NHLBI (ATP III)*	> 102cm	> 88cm		
Canada	Health Canada	> 102cm	> 88cm		
European	European Cardiovascular Societies	> 102cm	> 88cm		
Asian (including Japanese)	IDF	> 90cm	> 80cm		
Asian	WHO	> 90cm	> 80cm		
Japanese	Japanese Obesity Society	> 85cm	> 90cm		
China	Cooperative Task Strength	> 85cm	> 80cm		
Middle East, MEDITERRANE AN	IDF	> 94cm	> 80cm		
Sub-Saharan African	IDF	> 94cm	> 80cm		
Ethnic Central and South American	IDF	> 90cm	> 80cm		

syndrome *Recent AHA/NHLBI auidelines for metabolic an increased risk for CVD and diabetes at waist circumference thresholds of > 94 cm in men and > 80 cm in women and identify these as optional cut points for individuals or populations with increased insulin resistance.

IDF, International Diabetes Federation; AHA/NHLBI, American Heart Association/National Heart, Lung, and Blood Institute; WHO, World Health Organization. From Alberti KG, Eckel RH, Grundy SM, et al: Harmonizing the metabolic syndrome: a joint interim statement of the International Diabetes Federation Task Force on Epidemiology and Prevention; National Heart, Lung, and Blood Institute; American Heart Association; World Heart Federation; International Atherosclerosis Society; and International Association for the Study of Obesity. Circulation 120:1640, 2009.

atherosclerosis. 53 Free fatty acids, produced more readily in the visceral abdominal fat, may decrease insulin sensitivity, impair vascular reactivity, and also increase endothelial dysfunction. ⁵³

Low Birth Weight

The relationship between low birth weight and increased risk of obesity, hypertension, type 2 diabetes, stroke, and CVD later in life is well documented in epidemiological studies. 54 These associations remain strong even after adjustment for such lifestyle factors as smoking, physical activity, occupation, dietary habits, and childhood socioeconomic status. 55

A large number of studies have linked low birth weight to the later development of central adiposity. 55 A landmark cohort study of 300,000 men by Ravelli and coworkers ⁵⁶ showed that exposure to the Dutch famine of 1944-1945 during the first half of pregnancy resulted in low birth weight associated with significantly higher obesity rates at the age of 19 years. Subsequent research has confirmed the relationship between low

322 standard deviation of BMI. 80 In patients with class II and class III Munster (PROCAM) study, have reported a clear prognostic obesity, obstructive sleep apnea is common. 80 Lopez and inverse relationship between HDL-C levels and coronary artery colleagues 83 found the prevalence of obstructive sleep apnea to be disease morbidity and mortality, regardless of LDL-C levels. 99 greater than 70% in those with class II and class III obesity and more These outcomes reflect the need to identify ways to optimize the than 90% in patients with a BMI > 60.

A population-based prospective cohort study from 1989 2000 obesity, and dyslipidemia. found that a 10% weight gain predicted an approximate 32% increase in the apnea-hypopnea index and a sixfold rise in the risk Diabetes for development of moderate to severe obstructive sleep apnea. 84 The Sleep Heart Health Study showed a similar relationship between BMI and obstructive sleep apnea severity; the odds ratio for moderate to severe obstructive sleep apnea was 1.6 for each standard deviation increment in BMI. 85

Dyslipidemia

Atherogenic dyslipidemia is associated with an increased risk of adjusted percentage of obesity in adults with diabetes, an increase CVD, peripheral vascular disease, and stroke. 86.87 The condition is from 34.9% to 53.0%. 101 During that same period, the percentage of characterized by elevated triglycerides and low plasma levels of overweight or obese adults with diabetes increased from 70% to HDL-C, 88 often with elevated apolipoprotein B and non-HDL-C. It 83%. 102 is prevalent in patients with type 2 diabetes, metabolic syndrome, or established CVD. 89

Obesity heightens the risk of type 2 diabetes, hypertension, 19CVD, and dyslipidemia and reduces average life expectancy. 90,91 development of insulin resistance and type 2 diabetes. 12 In the characterized by high levels of triglycerides and LDL-C and low development of type 2 diabetes during a 16-year period. 103 levels of HDL-C, a combination that significantly increases the dyslipidemia, are interrelated, and each predicts CHD risk. 88

LDL particle number, and decreased HDL particle size. 12 These blacks, and 37% for whites. 104 lipoproteins can undergo a process of oxidation that results in the LDL. 93

metabolism of LDL-C and HDL-C and contributes to atherogenic CVD rates are lower. 105 potential. 93 Atherogenic dyslipidemia, characterized by elevated triglycerides and low levels of HDL-C, often with elevated The incidence of type 2 diabetes in this age group has increased in apolipoprotein B and non-HDL-C, is common in patients with tandem with obesity, rising by a factor of more than 10 in the past established CVD, type 2 diabetes, or metabolic syndrome and two decades. 106 This trend in children and adolescents is contributes to both macrovascular and microvascular residual risk. accelerating in both developed and developing countries. 107 As the

involving patients at different levels of risk shows that lipid and in glucose metabolism that can lead to type 2 diabetes. 107 lipid protein abnormalities are responsible for residual CVD risk in patients receiving statin therapy. 95 A recent meta analysis impaired glucose tolerance or impaired fasting glucose to diabetes including 90,056 subjects (18,686 with diabetes) from 14 is 25% during 3 to 5 years. 108 The only longitudinal study published randomized trials reported that for each millimole per liter thus far on the natural history of normal and impaired glucose decrease in LDL-C, statin therapy reduced the risk of major tolerance in children and adolescents showed that children with vascular events by 21%. Nevertheless, 14% of patients in the statin impaired glucose tolerance who had greater degrees of obesity at group suffered a cardiovascular event compared with 18% baseline and those who continued to gain weight rapidly randomized to placebo. 96,97

The importance of dyslipidemia as a major contributor to CVD risk is underlined by the INTERHEART study, 98 a global casecontrol trial in 52 countries, in which dyslipidemia was responsible for 54% of population attributable risk for myocardial infarction. Extensive evidence supports elevated triglycerides and low HDL-C levels as predictors for CVD, independent of LDL-C. 90 Observational trials, such as the Prospective Cardiovascular

management of patients with metabolic disorders, such as diabetes,

CVD affects millions of adults with diabetes and is a major cause of morbidity and mortality. Evidence indicates that the CVD burden among diabetics is increasing. Between 1997 and 2007, those aged 35 years or older with diabetes and a diagnosed CVD condition (ie, CHD, stroke, or other heart condition) increased from approximately 4 million to almost 6 million. 100 Between 1994 and 2007, there were also unfavorable upward trends in the age-

Obesity clearly increases the risk of developing type 2 diabetes; inflammatory adipokines may also play a role. 12 Large population studies have confirmed the links between excess weight and the Metabolic syndrome is closely associated with obesity and a Nurses' Health Study, which observed close to 85,000 female clustering of CVD risk factors, including a dys lipidemia nurses, a BMI > 25 was the single most important factor for the

A 20-year follow-up study of ethnicity, obesity, and risk of type relative risk of cardiovascular or cerebrovascular events. 92 2 diabetes in a Nurses' Health Study cohort of 78,419 apparently Triglycerides, HDL-C, and LDL-C, the components of atherogenic healthy women found that for each 5-unit increase in BMI, the multivariate relative risk of diabetes was 2.36 for Asians, 2.21 for Individuals with obesity and metabolic syndrome present with Hispanics, 1.96 for whites, and 1.55 for blacks. For each 5-kg weight increased concentrations of very-low-density lipoprotein (VLDL) gain between the age of 18 years and the year 1980, the risk of particles, increased triglycerides and small particle LDL, increased diabetes increased by 84% for Asians, 44% for Hispanics, 38% for

A study of 91,246 patients in 27 European countries investigated formation of foam cells and enhanced monocyte binding, which the impact of adiposity on the frequency of diabetes and CVD. Data leads to the early stages of atherosclerotic plaque. Small-particle showed that waist circumference predicted increased age- and LDL has greater atherogenic potential 93 and is more common in BMI-adjusted risks of CVD and diabetes. In women, odds ratios for individuals with diabetes. 94 As risk indicators, total CVD per 1 SD increase in waist circumference were 1.28 in cholesterol/HDL and LDL/HDL ratios have greater predictive northwest Europe, 1.26 in southern Europe, and 1.10 in eastern value than isolated parameters used independently, particularly Europe. Values for diabetes were 1.72, 1.45, and 1.59. Despite regional differences in cardiovascular risk factors and CVD rates, Elevated triglyceride concentrations are associated with greater abdominal obesity had a similar impact on the frequency of circulating numbers of triglyceride-rich VLDL particles and higher diabetes across Europe. The authors concluded that increasing levels of VLDL cholesterol, an environment that alters the abdominal obesity may offset future declines in CVD, even where

Diabetes is a serious weight-related condition in adolescents. prevalence of obesity increases, its health implications are Extensive evidence from large prospective clinical trials becoming more evident. The earliest alterations are abnormalities

> In adults, the likelihood of the progression of patients with developed type 2

diabetes. 109 Smaller studies of at-risk populations also suggest a people, a 10% weight loss was correlated with a 26% decrease 323 high likelihood of progression to type 2 diabetes. 109 Preliminary - in the apnea-hypopnea index, showing that even minimal weight data from Canada indicate that adolescents with type 2 diabetes will be at high risk for limb amputation, kidney failure requiring dialysis, and premature death. 106

The SEARCH for Diabetes in Youth study 110 found significant ethnic variations in the prevalence of type 2 diabetes in children decrease in apnea-hypopnea index from baseline. At 3-month aged 10 to 19 years. The disease accounted for only 6% of all follow-up, 61% of the patients in the intervention group were diabetic cases diagnosed in non-Hispanic whites compared with considered cured of sleep apnea compared with 32% in the control 22% of all diabetic cases diagnosed in Hispanics . In American Indians, type 2 diabetes has overtaken type 1 in prevalence among children, accounting for 72% of all cases of diabetes. Data for this study were obtained largely through chart review and may underestimate the prevalence of type 2 diabetes. As the ethnic diversity of the US population continues to increase, the epidemic of childhood obesity may make pediatric type 2 diabetes a growing that emphasizes the importance of treating both disorders. 80 public health concern. 13

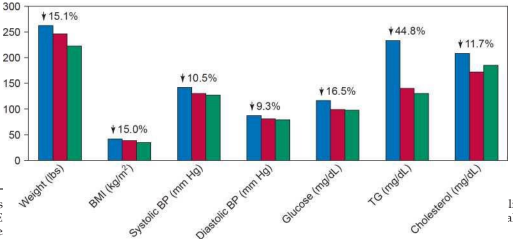
EFFECT OF WEIGHT LOSS ON OBESITY COMORBIDITIES 23 24

loss can be beneficial in patients with obstructive sleep apnea. 84 A randomized study of the effect of a low-calorie diet and supervised lifestyle counseling on sleep-disordered breathing produced a 40% group. Changes in apnea-hypopnea index were strongly correlated with changes in weight and waist circumference and were maintained at 1-year follow-up. 113 Data indicate that not only is obesity a risk factor for the development of obstructive sleep apnea, it may also be a consequence of obstructive sleep apnea, a finding

Previous meta-analyses of clinical trials on the effects of weight reduction on blood pressure show that weight loss is important in the prevention and treatment of hypertension. 114 In the Diabetes Prevention Program, a 2.8-year follow-up I found that weight loss was about 5.6 kg in the lifestyle man- I agement arm, 2.1 kg in the metformin arm, and 0.1 in the placebo group. There was a small significant decrease of 3.3 mm Hg in systolic blood pressure and a decline of 19 3.1 mm Hg in diastolic blood pressure in the lifestyle man agement group, suggesting that at least in the short term, weight loss was associated with some degree of decrease in blood pressure. 115-117

PRÉMIER, an NHLBI-sponsored multicenter randomized trial of 810 adults with prehypertension or stage 1 hypertension , compared the effect of advice only with established lifestyle interventions (eg, weight loss, dietary changes, and increased physical activity) for blood pressure control. The two intervention groups, established lifestyle interventions and established lifestyle interventions plus the DASH diet, reduced estimated 10-year CHD risk by 14% and 12%, respectively . 118 Other investigations have reported a linear association between changes in systolic blood pressure and weight, even with a small amount of weight loss. They have also documented that weight loss is the most important determinant of decreases in systolic blood pressure. 119,120

EFFECT OF MODERATE WEIGHT LOSS ON CARDIOMETABOLIC RISK



Losing weight has 111 The Look AHE overweight or obe 24diabetes, investi lifestyle interventi

lipidemia (Fig. 19-4). al conducted in 5145

ants in the intensive Percent changes are initial visit to final visit ent in cardiovascular

fitness. The intensive lifestyle intervention was associated with an increase from 46% to 73% of participants who met the American Diabetes Association (ADA) goal of A1c < 7% and a doubling in the aperquit age of individuals who met all three of the ADA goals for glycemic control, hypertension, and dyslipidemia. 112

Weight loss also interested district investeop caparage stream throughour inspiritors different also interested district through the population-based prospective cohort struct al flamact of weight loss on the metabolic syndrome. Diabetes Obes Metab 4:407, 2002.)

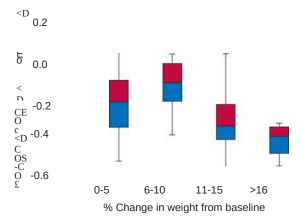


FIGURE 19-5 Box plots of percentage change in C-reactive protein (CRP) level from baseline over categories of percentage change in weight from baseline across all included weight loss interventions. (From Selvin E, Paynter NP Erlinger TP: The effect of weight loss on C-reactive protein: a systematic review. Arch Intern Med 167:31, 2007.)

Weight loss in overweight adolescents is also associated with a decrease in blood pressure ⁶⁷ as well as with reduced sensitivity of blood pressure to salt and other cardiovascular risk factors, such as dyslipidemia and insulin resistance. ¹²¹ In studies that reduce BMI by about 10%, short-term reductions in blood pressure were in the range of 8 to 12 mm Hg. An analysis of 33 weight loss interventions found that for each 1 kg of weight loss, the mean change in CRP level was - 0.13 mg/L, suggesting that weight loss is an effective non-pharmacological strategy for lowering of serum CRP levels (Fig. 19-5). ¹²² Although difficult, weight loss, if it is successful, is extremely effective. ^{67,121}

Despite these findings, application of data from observational trials must be carefully considered. Numerous studies have documented an obesity paradox in which overweight and obese individuals with established CVD (including hypertension, heart failure, CHD, and peripheral arterial disease) have a better prognosis compared to patients who are not overweight or obese. ³⁹ Conversely, patients with a healthy weight BMI (18.5 to 24.9 kg/m ²) and high body fat are at high risk for cardiometabolic dysregulation, metabolic syndrome, and CVD.

These findings, however, are controversial. ¹²³ Data assessing mortality based on body fat and lean mass rather than on BMI or weight alone have shown that subjects who lose body fat rather than lean mass have a lower mortality. ¹²⁴ Estimates for all-cause mortality, obesity-related causes of death, and other causes of death showed no statistically significant or systematic differences between BMI and other variables. ¹²⁵ Although an obesity paradox exists with the use of either baseline BMI or baseline percentage fat criteria, studies support the safety and potential long-term benefits of purposeful weight loss in overweight and obese patients with CHD. ^{39,126}

TREATMENT OPTIONS

Lifestyle Interventions: Diet, Behavioral Modification, and Exercise

A study examined the risk of CHD associated with excess weight in 42,351 men from the Health Professionals Follow up Study and 76,703 women from the Nurses' Health study. A total of 2771 incident cases of CHD among men and 2359 among women were documented during 16 years of follow-up. Overall, the relative risk of CHD associated with a BMI > 30 compared with a BMI of 18.5 to 22.9 was 2.13 among men and 2.48 among women. The risk of CHD increased with BMI, with and without hypercholesterolemia, hypertension, or diabetes. The authors estimated that more than a third of all CHD incidents in US men

and women may be attributed to excess weight. 127

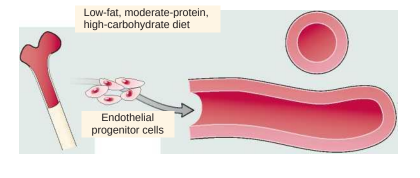
Sedentary lifestyles and poor physical fitness are major contributors to the current obesity and CVD pandemic. ¹²⁸ Thus, the main treatments of overweight and obesity include dietary changes, increases in physical activity, and other behavioral modifications. Technology-based approaches have also started to emerge. ¹²⁹ Whichever approach is used, regular activity and appropriate energy intake play critical roles in prevention and management of the negative health consequences of diabetes, obesity, and other cardiovascular diseases. ¹³⁰

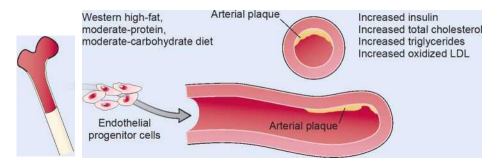
Nutrition remains a cornerstone of the effort to prevent CVD. Risk for heart disease can be minimized by adopting healthy eating patterns early in life and establishing life-long dietary habits of avoiding excess saturated fat, *trans*-fat, and salt. ¹³¹ Dietary changes can also improve CVD risk factors. For prevention of CVD and CVD risk factors, dietary choices that improve the overall quality of the diet are preferred to specific dietary components. The American Heart Association recommends that individuals consume a variety of fruits, vegetables , and grain products, especially whole grains. It also recommends fat-free and low-fat dairy products, legumes, poultry, and lean meats as well as fish, preferably oily fish, at least twice a week. ¹³² These foods should replace less nutrient dense ones to prevent weight gain associated with additional calorie consumption.

The "whole diet" approach greatly increases the odds of achieving a "prudent diet," characterized by a high intake of vegetables, fruits, legumes, fish, poultry, and whole grains. In a large, 18-year prospective study of 72,113 women, greater adherence to the prudent pattern was related to a lower risk of cardiovascular and total mortality. In contrast, greater adherence to the Western pattern (ie, high intake of red and processed meat, refined grains, French fries, and sweets and desserts) was linked to a higher risk of CVD, cancer, and total mortality. ¹³³

The optimal balance of macronutrients for weight loss and weight loss maintenance is a topic of much debate. A 2009 study found that macronutrient composition had a negligible effect on 2year weight loss outcomes in a cohort of overweight adults. 134 Instead, compliance with calorie restriction and counseling session attendance were strongly correlated with weight loss and, consequently, a reduction in risk for CVD and diabetes. Nevertheless, fat intake should be of particular concern in any lifestyle intervention. Consumption of diets rich in saturated fatty acids is highly correlated with metabolic syndrome and an increased expression of genes involved in inflammation processes in adipose tissue. 135,136 Whereas high-fat, high-protein, lowcarbohydrate diets have been associated with short-term improvement in LDL-C, HDL-C, and blood pressure, recent evidence suggests that this diet profile can elevate the risk of CVD without altering classic CVD risk factors. Instead, high-fat, highprotein diets can elevate circulating nonesterified fatty acids and suppress endothelial progenitor cell production, thereby increasing arterial plaque buildup (Fig. 19-6). 119,137

Insulin resistance due to atherogenic dyslipidemia in skeletal muscle may be the primary driving force in the development of the metabolic syndrome. ¹³⁸ Chronically ele vated serum free fatty acids could lead to further metabolic complications and morbidity. The National Cholesterol Education Program's Adult Treatment Panel III report recommen dations for diet composition for patients with metabolic





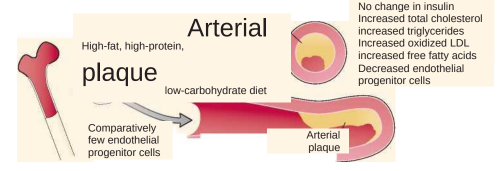


FIGURE 19-6 High-protein, low-carbohydrate diet and atherosclerosis. (From Smith SR: A look at the low-carbohydrate diet. N Engl J Med 361:2286, 2009.)

syndrome called for low intake of saturated fats, *trans* -fats, and cholesterol; reduced consumption of simple sugars; and increased intake of fruits, vegetables, and whole grains. ¹³⁹ Lowfat diets, such as the one used in the Women's Intervention - Nutrition Study, have been implemented in the long term to successfully reduce disease risk. ¹⁴⁰ In a cohort of patients from the Framingham Offspring Cohort, the consumption of a diet consistent with the principles of the Mediterranean-style diet was associated with protection against metabolic syndrome. ¹⁴¹

Data also show that replacement of saturated fats with omega-6 polyunsaturated fatty acids can lower risk for heart disease events by 24%. ¹⁴² An emphasis on monounsaturated fatty acids in the diet can lead to better inflammatory gene expression profiles, decreases in serum LDL-C concentrations, and an increase in plasma and adipose tissue oleic acid content. ¹³⁶ Frans -fatty acids adversely affect both LDL-C and HDL-C levels and increase the risk for CHD, even at low levels of dietary intake. ^{143,144} Elimination of exposure to *trans* -fatty acids could have a powerful population impact, potentially protecting 30,000 to 100,000 Americans from death related to heart disease. ^{143,144}

In general, comprehensive lifestyle modification programs

delivered in person induce a loss of approximately 10% of initial weight in 16 to 26 weeks of treatment. 145 Combined with energy restriction, dietary changes can increase weight loss, prevent weight regain, and reduce the risk of death related to heart disease. 146 The use of portion-controlled food, which typically involves meal replacement products, ²⁵ has been associated with medically significant weight loss. Energy restriction can, however, have negative consequences if fat-free mass is metabolized along with adipose tissue. Reductions in fat-free mass can reduce basal energy expenditure because of losses in metabolically active lean tissues. 147,148 Therefore, dietary programs that support fat-free mass maintenance and skeletal muscle biogenesis are advisable. Low-protein, calorie-restricted diets are associated with increased loss of fat-free mass. 149 Conversely, higher protein diets can preserve fat-free mass and improve blood lipid profiles. 150

Behavioral treatment is widely acknowledged as an essential component of effective lifestyle interventions. Traditional behavioral counseling models have assumed that patients and clients will change behavior simply by learning the facts about diet and exercise. There are, however,

significant limitations to this counseling strategy. ¹³¹ Additional emphasis is needed on ways to implement current guidelines in a contemporary society characterized by wide availability of unhealthy, energy-dense food and stressful lifestyles. In this sense, it may be more important to focus on barriers to implementation before providing specific nutrition counseling.

Self-monitoring, stimulus control, exercise, and cognitive restructuring represent four key components of behavioral modification. ¹⁵¹ Some evidence suggests that the ability to balance the immediate gratification of appetizing, energy-dense foods with their long-term consequences on weight and health may be dependent on the functional capacity of cognitive-emotional processing systems in the brain. ¹⁵² These neurocognitive resources may also be critical for maintenance of eating-related goals, cognitive control of eating behavior, and suppression of automatic biases toward food-related stimuli. ^{152,153}

Group sessions led by registered dietitians or behavioral psychologists are an effective venue for delivery of behavior therapy. These sessions also provide a combination of social support and friendly competition. Web-based programs represent a new frontier for behavioral treatment in obesity medicine as a low-cost and wide-reaching alternative to on-site treatment. Online programs could facilitate ongoing patient-provider contact, a key factor for long-term weight control. ¹⁵⁴

Regular physical activity is an essential component of primary prevention of CVD and obesity. ¹⁵⁵ Federal guidelines now recommend exercise for both prevention of disease and improvement of health. ¹⁵⁶ Physical activity and exercise both have positive effects on maintenance and promotion of healthy body weight. ¹⁴⁶ In the Look AHEAD trial, greater self-reported physical activity was the strongest correlate of weight loss after 1 year. ¹⁵⁷

Most public health guidelines recommend that adults participate in 30 minutes of moderate-intensity physical activity on most days of the week. 158 Aerobic endurance training is effective for improving maximum oxygen uptake and modifying cardiovascular risk factors associated with the development of coronary artery disease. 159 Resistance training, which has long been touted for its strength-enhancing effects, has recently been recognized for its relationship to health and disease risk. 159,160 Moderate- to high-intensity resistance training performed 2 or 3 days per week is associated with improvements in CVD risk factors in the absence of significant weight loss. 146,160,161 The addition of a muscle strengthening exercise program to a weight loss intervention may help conserve fat-free mass and basal energy expenditure and facilitate weight loss maintenance. 162 When it is paired with regular aerobic physical activity, resistance training may represent a feasible exercise intervention to promote healthy body composition and to prevent excess adiposity ¹⁶³ (Table 19-3). ¹⁶⁰ Even among those in the pre-obese range, vigorous physical activity can decrease the risk of heart failure. 164

Guidelines for glycemic control have become controversial . When levels are set too low (eg, hemoglobin A1c levels as low as 6.5% to 7%), they burden patients with complex treatment programs, hypoglycemia, weight gain, and costs. A review of large randomized trials in patients with type 2 diabetes suggests uncertain benefits and the need for a different approach—one that prioritizes well-being, healthy life styles, preventive care, and cardiovascular risk reduction. ¹⁶⁵ Important challenges also remain in preventing weight regain after weight loss interventions. ²⁵

Weight Loss Surgery

Bariatric surgery is associated with significant weight loss and improvement in diabetes, hypertension, and obstructive

TABLE 19—3 Comparison of Effects of Aerobic Endurance Training with Strength Training on Health and Fitness Variables

Variable	Aerobic Exercise	Resistance Exercise	
Body composition Bone mineral density Percentage body fat	TT 44	TT 4	
Lean body mass Muscle strength	0 0T	TT TTT	
Glucose metabolism			
Insulin response to glucose challenge Basal insulin levels Insulin sensitivity	44 4 TT	44 4 TT	
Plasma lipids and lipoproteins HDL-C	To	To	
LDL-C Triglycerides	40 44	40 40	
	4.4		
Resting heart rate Stroke volume, resting and maximal	44 TT	0 0	
Cardiac output, rest	0	0	
Cardiac output, maximal Systolic blood pressure at rest	TT 40	0 0	
Diastolic blood pressure at rest Vo₂max	40 TTT	0 T 0	
Submaximal and maximal endurance time	TTT	TT	
Submaximal exercise rate-pressure product Basal metabolic rate	444 To	44 T	

T, values increase; 4, values decrease; 0, values

unchanged; 1 arrow, small

effect; 2 arrows, moderate effects; 3 arrows, large effect; HDL-C, high-density lipoprotein cholesterol; LDL-C, low-density lipoprotein cholesterol Williams MA, Haskell WL, Ades PA, et al: Resistance exercise in individuals with and without cardiovascular disease: 2007 update: a scientific statement from the American Heart Association Council on Clinical Cardiology and Council on Nutrition, Physical Activity, and Metabolism. *Circulation* 116:572, 2007.

sleep apnea. 166 It remains the most diabetes. 167 A substantial body of literature shows that it achieves significant long-term weight loss with minimal mortality or complications. 168,169 Studies also indicate that weight loss surgery confers a survival advantage on those who undergo it compared with community controls. 170,171 Landmark findings from the Swedish Obese Subjects study show an estimated 28% reduction in the adjusted overall mortality rate compared with conventionally treated controls. 172 Similar outcomes have been cited in other reports. 168

Weight loss surgery reduces calorie intake by modifying the anatomy of the gastrointestinal tract through restriction, malabsorption, or a combination of the two techniques. ¹⁶⁸ Ensuing changes in the gut-brain axis alter peptides that may regulate appetite and satiety ¹⁷³ (e.g., ghrelin, glucagon-like peptide, and pancreaticpolypeptide). Amongseveral competing approaches for the management of severe obesity, the general trend is toward combined restrictive-malabsorptive procedures, such as gastric bypass and biliopancreatic diversion. ¹³⁸

In the United States, laparoscopic Roux-en-Y gastric bypass is the most effective procedure for weight loss and is considered the "gold standard" operation for long-term weight control. ¹⁶⁸ It accounts for more than 90% of all bariatric operations. Laparoscopic adjustable gastric banding is the second most commonly performed procedure. ¹⁶⁸ Data show that laparoscopic sleeve gastrectomy, which is becoming popular as a stand-alone operation for the treatment of severe obesity and related diseases, ¹⁷⁴ is safe and effective, with

From Aronne LJ, Wadden T, Isoldi KK, Woodworth KA: When prevention fails: obesity treatment strategies. Am J Med 122(Suppl 1):S24, 2009.

results similar to those of gastric bypass. ¹⁷⁵ Investigational treatments include intragastric balloon, gastric pacing, and endoluminal interventions.

A recent systematic review ¹⁶⁹ summarized the risks and benefits of weight loss surgery. Comorbidities in all groups improved after procedures. Statistically fewer people had metabolic syndrome, and there was higher remission of type 2 diabetes than in nonsurgical groups. In one large cohort study, the incidence of three of six comorbidities assessed 10 years after surgery was significantly reduced compared with conventional treatment. Mortality ranged from none to 10%. Major postoperative adverse events, some necessitating reoperation , included anastomosis leakage, pneumonia, pulmonary - embolism, band slippage, and band erosion.

Pharmacotherapy

In a large number of patients who are unable to reduce weight by nonpharmacological measures, drug therapy can help them reach weight control goals. However, drug treatment should be considered only part of a systematic weight management program that includes dietary and lifestyle changes. ¹⁷⁶ The Practical Guide: Identification, Evaluation, and Treatment of Overweight and Obesity in Adults recommends pharmaco therapy for patients with cardiometabolic risk and a BMI of 27 to 29.9 or those with a BMI > 30. ¹⁷⁷

Available pharmacological options for weight loss that are approved by the Food and Drug Administration (FDA) are limited to anorexiants (eg, phentermine); orlistat, a lipase inhibitor; and sibutramine, a drug that works by acting on appetite control centers in the brain (Table 19 -4). ^{25,178} Agents approved for other uses, such as antidepressants, have promoted weight loss in preliminary clinical trials. These include bupropion, a drug approved as an antidepressant and smoking cessation aid ¹⁷⁹; topiramate, a therapy approved for migraine prevention and the treatment of seizures ¹⁸⁰; and exenatide, a novel incretin mimetic approved by the FDA for adjunctive therapy to improve glycemic control in patients with type 2 diabetes. ¹⁸¹ Cetilistat, an oral, nonabsorbed synthetic lipase

inhibitor, has reached the phase III trial stage of development. 4,182 However, rimonabant, a cannabinoid 1 receptor antagonist that showed efficacy in producing clinically significant weight loss, has been withdrawn from the market because of safety concerns. 183

Emerging data from recently completed and ongoing randomized clinical trials suggest that certain combination drug therapies in development may have greater efficacy than currently available single-drug therapies in terms of weight loss and reduction in risk factors. ¹⁷⁹These drugs include Contrave, a combination of bupropion and the opioid antagonist naltrexone, ^{184,185} and pramlintide/metreleptin, a novel, integrated neurohormonal approach that combines amylin and leptin. ¹⁸⁶

CONCLUSION

Currently, one third of children and two thirds of adults are overweight or obese; this trend has persisted for the last two decades and shows no sign of abatement. ^{2,4} Obesity tracks from childhood into adulthood, with serious medical and economic consequences throughout the life course. ¹⁸⁷ One recent estimate suggests that if the current trend continues, obesity will account for more than \$860 billion or more than 16% of health care expenditures in the United States by 2030. ¹⁸⁸ Another projects that by 2035, the prevalence of CHD will increase by a range of 5% to 16%, with more than 100,000 excess cases of CHD attributable to increased adolescents obesity. ¹⁸⁹

To date, most public health strategies to address obesity have focused on efforts targeted at individuals, such as health education and behavioral skills training. However, these have turned out to be largely ineffective and unsustainable. ¹⁸⁷ Similarly, the majority of research on obesity prevention has ignored the combined effect of larger societal and political factors in perpetuating the obesity crisis. The lack of success



^{*}Data obtained from only one trail.

^{*}Data obtained from the Diabetes Prevention Program Trial; individuals with impaired fasting glucose were treated with drug

328 in stopping the spread of obesity underscores the need for a new approach that takes into account the sociopolitical, economic, and environmental forces that interact to create an obesogenic environment.

Individual choice is strongly influenced by the interaction of forces that take place at multiple socioenvironmental levels. ¹⁹⁰ These include interpersonal (family, peers, and social networks), community (schools, worksites, and institutions), governmental (local, state, and national policies), and biological processes (genes, molecular and cellular interactions, and the workings of major organ systems). ¹⁹⁰ Change at all of these levels is required to replace unhealthy social norms of diet and physical activity, to increase access to and availability of fresh fruits and vegetables, to provide opportunities for physical activity, and to promote a culture that values and encourages preventive care.

Reliance on free markets and free choice has failed to stem the obesity crisis. Investment in integrated, cross-disciplinary research is required, as are multilevel efforts to address social, economic, and environmental factors that create and sustain the conditions fostering obesity. A multilevel strategy rooted in a socioecological framework is consistent with the World Health Organization's work on social determinants of ¹⁹ health. ¹⁹² We already know what needs to be done. The question is whether we have the political will to change policies and practices that undermine public health.

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CHAPTER 20

Tobacco Use, Passive Smoking, and Cardiovascular Disease: Research and Smoking Cessation Interventions

Russell V. Luepker

KEY POINTS

- The smoking of tobacco products increases the risk of cardiovascular diseases, including myocardial infarction, stroke, and peripheral artery disease.
- The continued smoking of tobacco products in patients with cardiovascular disease increases the risk of recurrent events, including sudden death, and smoking cessation reduces this risk
- Environmental tobacco smoke (secondhand smoke) increases the risk for cardiovascular and other diseases in nonsmokers.
- Cigarette smoking rates among adults and youth have declined in the United States during several decades.
- Effective behavioral and pharmacologic methods are available to help smokers quit.
- Effective educational methods are available to prevent youth from starting to smoke.
- Policies and regulations including smoke-free spaces, limitations on advertising, enforcement of youth access laws, and increased cigarette excise taxes have contributed importantly to reductions in tobacco use.

There are approximately 43 million adult smokers in the United States. ¹ It is estimated that 443,000 deaths annually are attributable to cigarette smoking (Fig. 20-1). These deaths are replenished by an estimated 3,900 teenagers per day who start the smoking habit. ² The economic costs are estimated to be more than \$96 billion per year in medical expenses and \$97 billion in lost productivity. ³ Worldwide, tobacco is estimated to cause more than 5 million deaths per year. ⁴ The costs in lost health and in human suffering are incalculable.

The causal links between cigarette smoking and human disease are incontrovertible. Smoking is linked to major cardio vascular diseases including sudden death, acute myocardial infarction, peripheral artery disease, and stroke. 5 Cigarettes are also linked to many cancers and are the prime factor in lung cancer. Smoking is linked to acute and chronic pulmonary diseases including emphysema. There is growing evidence that environmental tobacco smoke, which results in exposure of nonsmokers, poses health risks to that group. Lung cancer, respiratory tract infections, and asthma attacks among those exposed to environmental tobacco smoke are well recognized. 6 More recent evidence finds that environmental tobacco smoke causes heart disease and sudden infant death syndrome. ⁶ These observations and others have resulted in a widespread call for prevention of tobacco uptake by teenagers, cessation among smoking adults, restriction of smoking in the environment. This chapter discusses the scientific evidence relating active and passive tobacco smoking to cardiovascular risk and the benefits of quitting. It also describes the trends in cigarette rette use. Finally, individual and population intervention strategies cessation among youth and adults discussed.

EFFECTS OF CIGARETTE SMOKING ON CARDIOVASCULAR DISEASES

There is a wealth of evidence in the past five

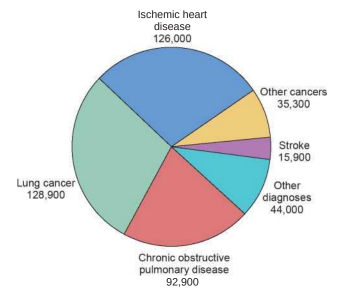
decades linking cigarette smoking to the major cardiovascular diseases, including myocardial infarction, sudden death, stroke, and peripheral vascular disease. ^{5,7} These associations are found across all age, gender, and ethnic groups.

The relationship of coronary heart disease (CHD) mortality to smoking status from the 1959-1965 Cancer Prevention Study is shown in Figure 20-2. CHD increases with age for both men and women; women have lower rates at all ages, and ever-smokers have significantly higher death rates than neversmokers. These differences are greatest in the younger age groups, in which the relative risk of CHD in smokers approaches 8. However, the differences remain into the older years, when relative risks are less but absolute risk is significantly greater. Smoking cessation among adults significantly reduces the risk of CHD and all cardiovascular diseases as shown in many populations. 8,9

Data from the Multiple Risk Factor Intervention Trial (MRFIT) of 316,099 white men also find a graded relationship between number of cigarettes and CHD death. ¹⁰ The relative risk for 1 to 25 cigarettes per day is 2.1, rising to 2.9 for daily smoking consumption above 25 cigarettes per day. Simi larly, MRFIT found that quitting smoking reduces cardiovascular disease mortality. ¹¹

In addition to age and gender effects, it is apparent that smoking-related disease also affects all the major ethnic and racial groups in the United States. ¹ The ill health effects of smoking cut across national boundaries as demonstrated in the Seven Countries Study of Keys and colleagues. ¹²

One of the most disturbing aspects of cigarette smoking is its strong association with sudden, unexpected death, particularly among younger individuals. Although sudden death is common among those with known cardiovascular disease, Esc obedo and Caspersen ¹³ found that only smoking predicted sudden death in those thought to be disease free. Similarly, in both men and women, acute myocardial infarction in younger individuals (younger than 50 years) is strongly associated with cigarette smoking. ¹⁴



'Average annual number of deaths, 2000-2004

FIGURE 20-1 About 443,000 U.S. deaths each year are attributable to cigarette smoking. (Source: cdc.gov.)

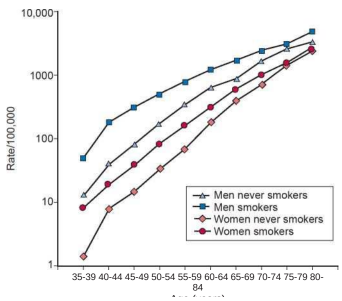


FIGURE 20-2 Coronary heart disease mortality. (Source: U.S. Department of Health and Human Services: Changes in cigarette-related disease risks and their implication for prevention and control, Monograph 8, Rockville, Md, 1997, U.S. Department of Health and Human Services, Public Health Services, National Institutes of Health, National Cancer Institute. NIH publication 97-4213.)

The interaction of cigarette smoking with other known risk factors is well studied. Some suggest that the effect is additive, whereas others find a multiplicative effect. Cigarette smoking adds to cardiovascular risk associated with lipids, obesity, diabetes mellitus, hypertension, oral contraceptive use, and electrocardiographic abnormalities. ¹⁵⁻¹⁷ The additive effects of smoking in relation to other major risk factors are seen in Figures 20-3 and 20-4 for 10- and 30-year follow- up of men and women aged 25 years at study entry. It is apparent that smoking adds significantly to risk. Smokers who continue in the habit after an acute myocardial infarction have significantly higher rates of recurrent events and death. ¹⁸ Individuals who quit smoking reduce that risk of a subsequent event. ¹⁸ Smokers also suffer increased rates of peripheral arterial disease with a relative risk of two to three times ¹⁹ and twice the risk of stroke. ²⁰

The mechanisms by which cigarettes affect cardiovascular diseases have been studied in both animal and human models. Both acute and chronic mechanisms are postulated, and it is likely that both contribute. There is accumulating evidence that smoking plays an important role in the basic atherosclerosis rhotic process. This is elegantly confirmed in the PDAY (Pathobiological Determinants of Atherosclerosis in Youth) study. 21 In this study, autopsies were performed on 1443 men and women aged 15 to 34 years who died of external causes, such as car accidents. Smoking was associated with an excess of fatty streaks and raised lesions in the abdominal aorta in these otherwise healthy individuals. 21 Injury of the arterial endothelium is suggested by some as the mechanism for the atherosclerotic lesions, but other mechanisms are also postulated. ²² Acute effects of smoking are demonstrated by the known short-term vascular effects of nicotine and the rapid improvement in prognosis with smoking cessation.

There are also known associations and interactions of smoking with other recognized elements in the causal chain and biomarkers for cardiovascular disease. ²³ For example, many studies find smoking to be inversely associated with high-density lipoprotein cholesterol, the "good cholesterol," whereas some report associations with elevated low-density and very-low-density lipoprotein cholesterol. Smoking has prothrombotic effects, including increasing platelet adhesion, increased fibrinogen, and decreased fibrinolytic activity. All are associated with increased clotting. ^{24,25} As the interest in atherosclerosis as an inflammatory disease grows, increased leukocyte counts and elevated C-reactive protein support an association. ²³

ENVIRONMENTAL TOBACCO SMOKE

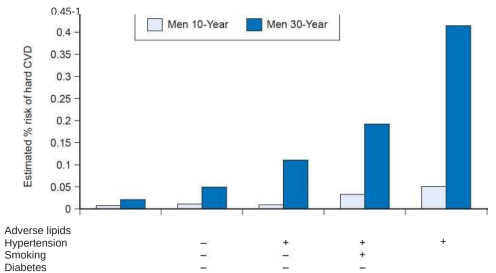
Recently, there is increased emphasis on exposure to environmental tobacco smoke among nonsmokers in the home, workplace, and public settings. This is more hotly debated than individual smoking because it affects those who may not have a choice about being exposed to tobacco smoke. In the 2006 Surgeon General's report *The Health Consequences of Involuntary Exposure to Tobacco Smoke*, cardiovascular disease was clearly implicated along with the demonstrated cancer and respiratory disease effects of environmental tobacco smoke. ⁶

A meta-analysis incorporating more home-based studies along with workplace studies (total of 1699 cases) showed an overall increased risk associated with passive smoking (RR = 1.49; 95% CI, 1.29-1.72) and suggested relative risks from workplace exposure similar to those from home-based exposure. ²⁶ In addition, environmental tobacco smoke is associated with sudden infant death syndrome and respiratory diseases in exposed children. ²⁷

The mechanism by which environmental tobacco smoke affects individuals is still debated, but considerable data are available. It is clear that mainstream smoke, which is inhaled by the smoker, differs from sidestream smoke, which is released into the environment immediately. ^{6.26} Sidestream smoke may be more toxic. It is apparent that nonsmokers who are exposed regularly to cigarette smoke develop a number of physiological changes. Some studies find that nonsmokers are more sensitive to these changes than those regularly exposed. ²⁸ These include the chronic effects of cigarette smoke, such as lower high-density lipoprotein cholesterol, increased fibrinogen, and platelet abnormalities. ⁶ It is also apparent that exposed nonsmokers have acute effects including endothelial dysfunction and lower exercise tolerance. ^{29,30}

These observations are compatible with pathological cardio vascular effects in nonsmokers exposed to the environment

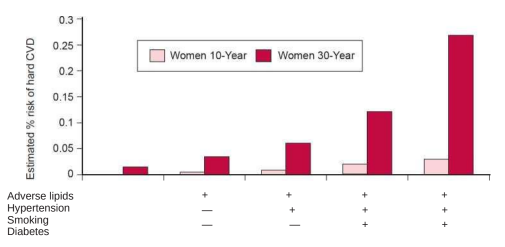




No risk factors profile: Total cholesterol=150 mg/dL; HDL cholesterol=60 mg/dL; untreated SBP=110 mm Hg; nonsmoker; nondiabetic. Adverse lipids: total cholesterol=260 mg/dL; HDL cholesterol=35 mg/dL. Hypertension: SBP=160 mm Hg, untreated.



FIGURE 20-3 The 10-year versus 30-year risk of hard CVD events for 25-year-old men with different risk profiles. (From Pencina MJ, D'Agostino RB Sr, Larson MG, et al: Predicting the 30-year risk of cardiovascular disease: the Framingham Heart Study. Circulation 119:3078, 2009.)



No risk factors profile: Total cholesterol=150 mg/dL; HDL cholesterol=60 mg/dL; untreated SBP=110 mm Hg; nonsmoker; nondiabetic. Adverse lipids: total cholesterol=260 mg/dL; HDL cholesterol=35 mg/dL. Hypertension: SBP=160 mm Hg, untreated.

FIGURE 20-4 The 10-year versus 30-year risk of hard CVD events for 25-year old women with different risk profiles. (From Pencina MJ, DAgostino RB Sr, Larson MG, et al: Predicting the 30-year risk of cardiovascular disease: the Framingham Heart Study. Circulation 119:3078, 2009.)

tobacco smoke. Recent estimates suggest that 23,000 to 70,000 deaths per year from acute myocardial infarction are associated with environmental tobacco smoke exposure. ³¹ This effect is much larger than that observed for lung cancer. These observations underlie the recent efforts and successes in reducing environmental exposures or secondhand smoke.

PREVALENCE AND TRENDS IN CIGARETTE SMOKING AMONG YOUTH

Cigarette smoking among youth is described in the 1994 Surgeon General's report, 32 and methods of prevention are

expanded in the 2000 Surgeon General's report. 33 Most smokers start this habit in their teenage years. It begins with social pressure based on friends, siblings, and parents who smoke. Youth believe that smoking makes you look more adult and is associated with social success, independence, rebelliousness, common themes in the teenage years. The environment provides important support through advertising, which reinforces the "coolness" of smoking. The highly effective Joe Camel ads and other media approaches were clearly aimed at and successful with new-onset smokers. 34 The basis of many tobacco lawsuits originated in this market practice. The most vulnerable period is in the sixth to eighth grades (ages 12 to 14 years), when much smoking initiation occurs.

TABLE 20—1 Percentage of High-School Students Who Reported Lifetime Cigarette Use,******** Current Cigarette Use/ and Current Frequent Cigarette Use - Youth Risk Behavior Survey, United States, 1991-2007 § 1993 (95% CI) 1995 (95% CI) 1997 (95% CI) 1999 (95% CI) 2001 (95% CI) 2003 (95% CI) 2005 (95% CI) 2007 (95% CI) CI) Cigarette Use Lifetime 11 70.1 70.4 63.9 54.3 (67.8-72.3)(68.1-70.8) (69.5-73.0) (68.2-72.1) (67.3-73.3)(61.6-66.0) (55.1-61.6) (51.2-57.3) (47.2-53.5)Current / 34.8 34.8 28.5 23.0 (24.8-30.3)(28.6 - 32.4)(32.5-37.2)(34.1-38.7)(32.3-37.4)(26.4-30.6)(19.8-24.2)(20.7-25.5)(17.6-22.6)Current 12.7 13.8 16.1 16.8 13.8 9.7 often 11 (10.6-15.3) (12.1-15.5) (14.8-18.7) (14.3-19.6) (12.3-15.5) (8.3-11.3) (7.9-11.0) (6.7-9.8)(13.6-19.1)

TABLE 20—2 Percentage of High-School Students Who Reported Current Cigarette Use, * by Sex, Race/Ethnicity, and Grade— Youth Risk Behavior Survey, United States, 1991-2007

Characteristic	1991 (95% CI)	1993 (95% CI)	1995 (95% CI)	1997 (95% CI)	1999 (95% CI)	2001 (95% CI)	2003 (95% CI)	2005 (95% CI)	2007 (95% CI)
Sex									
Female ·	27.3	31.2	34.3	34.7	34.9	27.7	21.9	23.0	18.7
Male ·	(23.9-31.0) 27.6 (24.6-30.9)	(29.1-33.4) 29.8 (27.4-32.3)	(31.0-37.7) 35.4 (32.9-37.9)	(31.8-37.6) 37.7 (35.0-40.6)	(32.3-37.7) 34.7 (31.8-37.7)	(25.6-30.0) 29.2 (26.7-32.0)	(19.2-24.9) 21.8 (19.8-24.1)	(20.4-25.8) 22.9 (20.7-25.3)	(16.5-21.1) 21.3 (18.3-24.6)
Race/Ethnicity §									
White, non-Hispanic ·	30.9	33.7	38.3	39.7	38.6	31.9	24.9	25.9	23.2
Female ·	(27.6-34.5) 31.7	(31.4-36.0) 35.3	(35.6-41.1) 39.8	(37.3-42.2) 39.9	(35.5-41.9) 39.1	(29.6-34.4) 31.2	(22.4-27.5) 26.6	(22.9-29.2) 27.0	(20.4-26.2) 22.5
Male ·	(27.1-36.7) 30.2 (26.5-34.3)	(32.6-38.0) 32.2 (29.4-35.0)	(36.3-43.5) 37.0 (33.7-40.5)	(36.6-43.2) 39.6 (35.8-43.5)	(35.4-42.9) 38.2 (34.6-41.8)	(28.7-33.7) 32.7 (29.7-35.9)	(22.9-30.5) 23.3 (20.7-26.0)	(23.4-31.0) 24.9 (22.2-27.7)	(19.6-25.7) 23.8 (20.2-27.8)
Black, non-Hispanic II	12.6	15.4	19.1	22.7	19.7	14.7	15.1	12.9	11.6
Female ^	(10.2-15.5) 11.3	(12.9-18.2) 14.4	(16.1-22.6) 12.2	(19.0-26.8) 17.4	(15.8-24.3) 17.7	(12.0-17.9) 13.3	(12.4-18.2) 10.8	(11.1-14.8) 11.9	(9.5-14.1) 8.4
Male ¹¹	(9.2-13.9) 14.1 (10.1-19.4)	(11.9-17.4) 16.3 (12.4-21.1)	(9.3-15.7) 27.8 (22.5-33.9)	(13.8-21.7) 28.2 (23.0-34.1)	(14.4-21.7) 21.8 (15.4-29.9)	(10.1-17.2) 16.3 (13.2-19.8)	(8.2-14.2) 19.3 (15.8-23.5)	(10.2-13.8) 14.0 (11.5-16.9)	(6.6-10.6) 14.9 (11.7-18.8)
Hispanic ·	25.3	28.7	34.0	34.0 (31.3-36.9)	32.7	26.6	18.4	22.0	16.7
Female ·	(22.5-28.2) 22.9	(25.8-31.8) 27.3	(28.7-39.6) 32.9	32.3	(29.0-38.6) 31.5	(22.4-31.2) 26.0	(16.1-20.9) 17.7	(18.7-25.8) 19.2	(13.5-20.4) 14.6
Male ·	(19.2-27.1) 27.8 (24.3-31.8)	(23.5-31.5) 30.2 (26.7-33.8)	(27.4-39.0) 34.9 (26.6-44.3)	(28.6-36.2) 35.5 (31.9-39.2)	(26.8-36.5) 34.0 (29.7-38.7)	(22.3-30.0) 27.2 (20.6-35.0)	(15.6-19.9) 19.1 (15.8-23.0)	(16.4-22.5) 24.8 (20.0-30.4)	(11.3-18.8) 18.7 (15.0-23.2)

Early surveys of national trends showed a steady rise in cigarette smoking among youth from 1968 to 1974. ³⁵ During that time, smoking of teenage girls began to exceed that of boys. However, from 1991 to 2007, a national sample of high school students found a steady fall of smoking rates among this group until 2003, when it leveled off (Table 20-1). ³⁶ Importantly , the category never-smoked was increasing. There were also race and

ethnicity differences, with non-Hispanic whites more likely to smoke than blacks or Hispanics in 2007 (Table 20-2). ³⁶

If the long-term health of the nation is to be improved, prevention of cigarette smoking initiation among youth is an essential element.

^{*}Ever tried cigarette smoking, even one or two puffs.

[·] Smoked cigarettes on at least 1 day during the 30 days before the survey.

[·]Smoked cigarettes on 20 or more days during the 30 days before the survey.

[§] Linear, quadratic, and cubic trend analyzes were conducted with a logistic regression model controlling for sex, race/ethnicity, and grade. These prevalence estimates are not standardized by demographic variables.

[&]quot;Significant linear and quadratic effects only (P < 0.06).

 $^{^{\}wedge}$ Significant linear, quadratic, and cubic effects (P < 0.05).

^{*******}Smoked cigarettes on at least 1 day during the 30 days before the survey.

[^]Linear, quadratic, and cubic trend analyzes were conducted with a logistic regression model controlling for sex, race/ethnicity, and grade in school. These prevalence estimates are not standardized by demographic variables.

[^] Significant linear, quadratic, and cubic effects (P < 0.05).

[^] Numbers for other racial/ethnic groups were too small for meaningful analysis.

[&]quot;Significant quadratic and cubic effects only (P < 0.05).

 $^{^{\}wedge}$ Significant linear and quadratic effects only (P < 0.05).

PREVALENCE AND TRENDS AMONG ADULTS

Cigarette consumption per capita for individuals aged 18 years and older rose steadily from 1900 to the late 1960s.

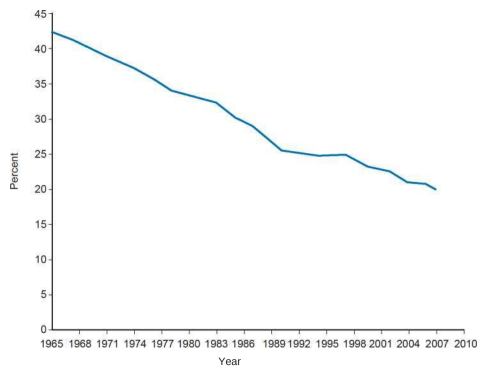


FIGURE 20-5 Trends in current cigarette smoking among adults, United States, 1965-2007. (Source: cdc.gov.)

Since that time, it steadily declined through 2007. ³⁷ This pattern is the result of complementary trends including increased levels of smoking cessation and increasing rates of never-smokers. The National Health Interview Survey found that 42.4% of adults smoked in 1965 and 19.8% in 2007 (Fig. 20-5). The proportion of never-smokers in 2008 was 57.6%, with former smokers (21.5%) exceeding current smokers (20.8%) (Fig. 20-6). These national rates confirm a declining number of adult smokers during the past 40 years. Although national rates have decreased, there is still significant variation by region of the country. ³⁸ In 2004, the percentage of adults who smoked ranged from a high of 28.3% in Ken tucky to 11.7% in Utah, with an average of 20%.

Whereas cigarette smoking among adults has substantially declined, a sizable portion of the population is still addicted. ³⁷ One of the national health objectives for 2010 is to reduce the prevalence of cigarette smoking among adults to 12% or less. With smoking rates of 19.8% in 2007, that goal may not be met. Prevalence was lowest among Asians (9.6%) and Hispanics (13.3%) and highest among American Indians/Alaska Natives (36.4%). Non-Hispanic whites (21.4%) and blacks (19.8%) were similar. Adults who lived below the poverty level were more likely to smoke (28.8%) than were those at or above this level (20.3%). Persons with graduate or professional post-college degrees were far less likely to smoke (6.2%) than were those with less than a high-school education (33%) or a GED diploma (44%). Smoking prevalence was far lower in those 65 years and older (8.3%) than in the younger age groups.

PREVENTION AND INTERVENTION AMONG YOUTH

School-Based Prevention Programs

Much of the effort in preventing smoking among youth has focused on the school setting. School-based prevention programs have generally been targeted at junior high

or middle-school children, when the habit begins. Unfortunately , school-based programming alone may have limited impact in the absence of active parental and community involvement. ³² At

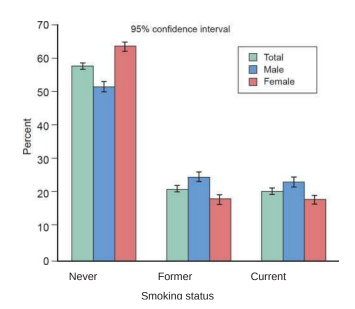


FIGURE 20-6 Percentage distribution of smoking status among adults aged 18 years and older, by sex: United States, January-June 2008. Note: Current smokers were defined as those who smoked more than 100 cigarettes in their lifetime and now smoke every day or some days. The analyses excluded 165 persons (1.3%) with unknown smoking status. (Based on data collected from January through June in the Sample Adult Core Component of the 2008 National Health Interview Survey. Data are based on household interviews of a sample of the civilian noninstitutionalized population.)

one time, it was assumed that simply educating youth about the harmful effects of smoking would be sufficient to prevent them from initiating cigarette use. ³⁹ However, it became apparent that information alone was not sufficient to deter adolescents from starting to smoke.

Social influence approaches have identified the social environment as a critical determinant of smoking onset. Rather than focusing on long-term disease risk, these interventions have stressed more immediate consequences of smoking, including negative social consequences. Adolescents are seen as often lacking in skills needed to resist peer pressure and other influences that promote smoking. ⁴⁰ As described in the 1994 Surgeon General's report, the main messages of successful skills-based interventions focus on the negative short-term social consequences of smoking, on the techniques of tobacco advertising that may be falsely appealing to adolescents, and on the socially salient advantages of being a nonsmoker.

Meta-analyses of school-based smoking prevention programs have indicated that these programs do have an impact on preventing smoking onset. ⁴⁰⁻⁴⁴ Furthermore, social influence approaches appear to be the most effective type of school-based program. On the basis of the results of a meta analysis, Rooney concluded that the best results were obtained by social influence programs that were delivered to sixth grade students, included booster sessions, concentrated the program within a short time, and used a peer to present the ^{44th} program.

Glynn ⁴⁵ listed essential elements of effective smoking - prevention programs based on a consensus panel. These - elements were summarized in the 1994 report of the Surgeon General ³² as follows:

- 1. Classroom sessions should be delivered at least five times per year in each of two years in the sixth through eighth grades.
- 2. Programs should emphasize the social factors that influence smoking onset, short-term consequences, and refusal skills.
- 3. Programs should be incorporated into the existing school curriculum.
- 4. Programs should be introduced during the transition from elementary school to junior high or middle school (sixth or seventh grade).
- 5. Students should be involved in the presentation and delivery of the program (peer teaching).
- 6. Parental involvement should be sought.
- 7. Teachers should be adequately trained.
- 8. Programs should be socially and culturally acceptable to each community.

Although some of these points might appear self-evident (eg, adequate training of teachers, social and cultural acceptability of programs), they are often overlooked in practice. Furthermore, high-risk populations, including those of low socioeconomic status and some ethnic minorities, present unique challenges as described by Sherman and Primack. ⁴⁶ These groups require additional attention.

Muller-Riemenschneider and colleagues ⁴⁷ evaluated the long-term effectiveness of smoking prevention programs published from 2001 to 2006. In 35 studies with a follow-up period of 1 to 10 years, they found that most studies reported long-term beneficial effects. They noted that school-based programs alone were least effective but that the addition of community-based and multisectoral approaches to school programs was most effective.

Community-Based Prevention Programs

Several model prevention programs have actively involved parents and the larger community in addition to schools. One example is described by Perry and colleagues ⁴⁸ in the context of the Minnesota Heart Health Program, a research and demonstration project designed to reduce cardiovascular disease at the community level. ⁴⁹ Perry and colleagues hypothesized that school-based smoking prevention would be more effective in communities in which multiple complementary school and community programs were established. ⁴⁸ Students participated in 5 years of school-based health education including peer-led prevention in a context in which adults were actively involved in community smoking cessation programs, and smoking restrictions were implemented both in schools and in the larger

community. Results were very encouraging, with smoking initiation significantly lower among students in the intervention community compared with students in the control community. These differences persisted throughout junior and senior high school. At the end of the twelfth grade, students in the intervention community evidenced 40% lower smoking prevalence than did students in the comparison community; 14.6% of students were weekly smokers at the end of high school compared with 24.1% in the reference community.

Other successful community interventions affecting youth include parental involvement, smoking restrictions (eg, schools, restaurants), and limitation of access to cigarettes through legal restrictions and increased taxes. ⁵⁰

State and Federal Prevention Initiatives

After the successful tobacco lawsuits and increases in cigarette taxes, many states operated comprehensive anti-tobacco programs. A major emphasis of these programs is on prevention of tobacco use in youth. Many of these initiatives include anti-tobacco media campaigns. ⁵¹ Florida launched an aggressive - media campaign that directly targeted the tobacco industry. ⁵² After 2 years, current cigarette use dropped from 18.5% to 11.1% among middle-school students and from 27.4% to 22.6% among high-school students. ⁵³

Flynn and coworkers ⁵⁴ found that a combination of media intervention and school-based programming fared better than school-based intervention alone. The media campaign included both radio and television spots that were broadcast as paid advertisements over local media. Reported smoking in the past week was 35% less among youth in the school-and-media condition than in the school-only condition (12.8% versus 19.8%).

Restrictions on smoking in public places including schools and on tobacco availability to minors may also reduce smoking prevalence, although findings are inconsistent. ⁵⁵ All 50 states and the District of Columbia adopted a minimum age of 18 years for the purchase of tobacco, but these laws are variably enforced.

Increased taxation also has an impact on adolescent smoking. Adolescents are more price sensitive than adults are. ⁵⁶ The impact of significantly higher prices may be even greater in discouraging initiation than in reducing consumption among existing smokers. Furthermore, revenue from these taxes can be dedicated to comprehensive tobacco control programs that will further reduce the onset of smoking in youth.

The Centers for Disease Control published *Best Practices for Comprehensive Tobacco Control Programs* in 1999 ⁵⁷ and updated this report in 2007. ⁵⁸ This document provides recommendations to the states for establishment of initiatives that include local community programs, chronic disease programs to reduce the burden of tobacco-related diseases, school interventions, enforcement, statewide programs, countermarketing, cessation methods, surveillance and evaluation , and administration and management. Recommended funding levels are given for each of these components. Local community programs can engage young people; school-based interventions can be linked with local community coalitions and statewide counteradvertising; enforcement can reduce access of minors; and statewide initiatives can provide skill, resources, and information for coordinated strategic implementation of community programs.

School-based prevention programs

Social influence approaches show good results.

Community-based prevention programs (may also enhance effects of schoolbased programs)

State and federal prevention initiatives

Anti-tobacco media campaigns (may enhance school-based programs) Restrictions on tobacco advertising

Restrictions on tobacco availability to minors

Restrictions on smoking in public places including schools Increased taxation (adolescents may be sensitive to price increases)

Limited work in adolescent cessation School-based cessation approaches Adolescent cessation in managed care

Cessation Among Youth

Most of the work done with youth has focused on prevention rather than on cessation of smoking. Many of the interventions demonstrated to be effective among adults have not achieved comparable success in adolescent populations.

A review examined the literature on smoking cessation in adolescents. 59 The authors evaluated 34 cessation studies. Program content for these studies was derived from a wide range of theoretical perspectives. Most studies were conducted in school settings that reach less than half of the potential population of adolescent smokers. End of treatment quit rates reported in 12 of the 17 studies averaged 20.7% (range, 0% to 36%); abstinence at follow-up declined to 13%. The most successful programs generally include some type of cognitivebehavioral intervention, such as instruction in coping skills and a focus on the immediate consequences of quitting.

A recently released handbook on cessation for youth, I Quit! What to Do When You're Sick of Smoking, Chewing, or Dipping, from the Centers for Disease Control and Prevention, provides helpful tips on quitting tobacco in this group. 60 Strategies for youth are summarized in Box 20-1.

CESSATION INTERVENTIONS AMONG ADULTS

Interventions with adults have focused on cessation rather than on prevention for obvious reasons, but there is a need for prevention in certain groups. Initiation among young adults is relatively common in certain settings, such as in the military. Klesges and colleagues 61 reported that 7% of never-smokers who entered the Air Force and completed basic military training were regular smokers 1 year later. Ethnic differences have also been found in smoking initiation; African Americans tend to initiate later than European Americans. 1 In some other countries, initiation of smoking tends to occur considerably later than in the United States. In China, for example, one cohort study found the mean age for starting smoking to be 22 years. 62 Until recently, it had been widely assumed that those who reached the age of 18 without cigarettes were very unlikely to initiate smoking as adults. However, the tobacco industry has increasingly targeted advertising and promotion efforts to young adults. In contrast to older adults, smoking prevalence has not been declining in the young adult (18- to 24-year-old) population.

Most of the published work with adults has been limited to individual smokers who have sought assistance in quitting. Unfortunately, individual smokers who are ready to quit and who seek help represent a very small proportion of the overall smoking population. 63 Furthermore, even among these smokers, 339 absolute long-term outcomes with formal quit smoking programs are disappointing. Intensive multisession group clinics generally produce no more than about 25% abstinence at 1 year. 64 Unaided quit attempts fare substantially less well; of the approximately 17 million smokers in the United States who attempt to quit on their own each year, less than 10% are successful.

Traditionally, interventions have focused on a single assisted quit attempt. A more effective approach may be to view smoking as a chronic disease and to support multiple quit attempts as necessary. 65 More recently, larger scale public health efforts address smoking cessation at the community level and target adults who may not volunteer for treatment or who may not be immediately interested in quitting. For many years, smoking and other tobacco use were seen as essentially learned behaviors. More recently, however, smoking and other tobacco products have been recognized as physically addictive. 66

Interventions Targeted at Individuals

There has been far more progress in developing effective smoking cessation interventions with adults than with adolescents. A recent clinical practice guideline panel of public 20 and private agencies sponsored by the US Public Health Service reviewed the literature and updated a 2000 report in 2008. 67 An overall conclusion of the panel was that effective smoking cessation treatments are available for adults. Recommendations from this panel are shown in Box 20-2.

Behavioral Treatments

Despite the addictive properties of nicotine, behavioral aspects of smoking are still seen as critical cessation tools. The most effective intervention programs, including those using medications, have included behavioral treatment components. A number of specific behavioral components are aversive smoking, intratreatment social support, problem solving/skills training, quit day, extratreatment social support, motivation, weight and exercise/fitness, diet/nutrition, contingency relaxation/breathing, and cigarette fading. Most of these specific treatment components have not been proven effective in isolation but may contribute to an overall multicomponent intervention.

Although aversive smoking approaches are effective (and indeed achieve the highest absolute abstinence levels), aversive techniques have largely gone out of favor. These techniques have included rapid smoking 68,69 and oversmoking or satiation. 70 Rapid smoking requires smokers to take very frequent puffs, typically every 6 seconds, for as long as they can tolerate the procedure. Oversmoking requires subjects to dramatically increase (perhaps double) their usual cigarette consumption for an arbitrary period, typically 1 week. Concerns have been expressed about the safety of rapid smoking and oversmoking, particularly in those with prevalent disease. 71 Acceptability of these techniques to smokers has been an additional issue. Other options have included reduced aversion techniques, 72 such as focused smoking (smoking at a regulated but slower rate) and smoke holding (retaining smoke in the mouth and throat while breathing through the nose).

Contingency Contracting

Several studies have required participants to submit monetary deposits that are refunded contingent on maintained



3OX 20-2 Smoking Cessation Clinical Guideline Recommendations for Adults ⁶⁶

- 1. Tobacco dependence is a chronic disease that often requires repeated intervention and multiple attempts to quit. Effective treatments exist, however, that can significantly increase rates of long-term abstinence.
- It is essential that clinicians and health care delivery systems consistently identify and document tobacco use status and treat every tobacco user seen in a health care setting.
- Tobacco addiction treatments are effective across a broad range of populations. Clinicians should encourage every patient willing to make a quit attempt to use the counseling treatments and medication recommended.
- Brief tobacco addiction treatment is effective. Clinicians should offer every patient who uses tobacco at least the brief treatments shown to be effective.
- 5. Individual, group, and telephone counseling are effective, and their effectiveness increases with treatment intensity. Two components of counseling are especially effective, and clinicians should use these when counseling patients making a quit attempt:
 - a. Practical counseling (problem solving/skills training)
 - b. Social support delivered as part of treatment
- 6. Numerous effective medications are available for tobacco dependence, and clinicians should encourage their use by all patients attempting to quit smoking, except when medically contraindicated or with specific populations for which there is insufficient evidence of effectiveness (ie, pregnant women, smokeless tobacco users, light smokers, and adolescents).
 - a. Seven first-line medications (five nicotine and two non- nicotine) reliably increase long-term smoking abstinence rates: bupropion SR, nicotine gum, nicotine inhaler, nico tine lozenge, nicotine nasal spray, nicotine patch, and varenicline.
 - Clinicians should also consider the use of certain combinations of medications identified as effective.
- 7. Counseling and medication are effective when used by themselves for treating tobacco addiction. The combination of counseling and medication, however, is more effective than either alone. Thus, clinicians should encourage all individuals making a quit attempt to use both counseling and medication.
- Telephone quitline counseling is effective with diverse populations and has broad reach. Therefore, clinicians and health care delivery systems should both ensure patient access to quitlines and promote quitline use.
- If a tobacco user is currently unwilling to make a quit attempt, clinicians should use the motivational treatments shown in the guideline to be effective in increasing future quit attempts.
- 10. Tobacco dependence treatments are both clinically effective and highly cost-effective relative to interventions for other clinical disorders. Providing coverage for these treatments increases quit rates. Insurers and purchasers should ensure that all insurance plans include the counseling and medication identified as effective as covered benefits.

abstinence ^{73,74} Contracts may also call for self-administered rewards for progressively longer periods of abstinence. Typically , contingency contracting has been included as part of multicomponent behavioral programs.

Social Support

Supportive intervention during direct contact with a clinician or in a group (intratreatment social support) increases smoking cessation rates. ⁶⁷ However, although social support from friends and family is strongly related to successful outcomes in smoking cessation treatments, efforts to systematically enhance natural social support as part of treatment intervention have generally proven unsuccessful. ⁷⁵

Relaxation Techniques

Progressive relaxation and deep breathing strategies have been employed for smoking cessation, although rarely in isolation. A major rationale for the use of these procedures is that smoking relapses are very likely to occur during negative emotional states. ⁷⁶ Relaxation training allows an alternative response for coping with negative emotions or stressful situations and with

the stress of quitting smoking and nicotine withdrawal effects. However, there is little evidence to support the effectiveness of relaxation training as a stand-alone technique. ⁷⁷

Coping Skills

Favorable results have been found for specific training in coping skills. Coping skills include problem solving and methods for management of stress and prevention of relapse. Shiffman ⁷⁸ found that a combination of cognitive (eg, mentally reviewing benefits of quitting) and behavioral (eg, physical activity, leaving a tempting situation) coping responses provided maximum protection against smoking in a potential crisis situation.

Reduced Smoking and Nicotine Fading

Nicotine fading is a nonaversive preparation technique based on the logical premise that withdrawal discomfort might be ameliorated if nicotine consumption is progressively reduced before abstinence. This premise may appear to be in conflict with the results of gradual reduction or cigarette "tapering" procedures. Strategies that have emphasized cutting down the number of cigarettes smoked have been almost uniformly unsuccessful. 79 Smokers typically appear to reach a "stuck point," often at 10 to 12 cigarettes per day. 79 For the typical smoker of approximately a pack per day, compensatory changes in puffing can compensate for reduced numbers of cigarettes at this level. Nicotine fading is an alternative in which smokers switch in a series of progressive steps during several weeks to cigarettes rated lower in tar and nicotine 80 or use commercially available nicotine reduction filters. 81 These procedures have not proven successful in improving smoking cessation outcomes.

Multicomponent Treatment Programs

The most successful behavioral programs have incorporated multiple treatment components. Emphasis has been placed on both initial preparation for quitting and longer term main -tenancy. Reported long-term abstinence rates for these multi-component treatment programs have approached 50%. 82

Hypnosis and Acupuncture

There are numerous approaches to smoking cessation that are not primarily behavioral or pharmacological. Two commonly advertised methods are hypnosis and acupuncture. Unfortunately , there are few good studies of these methods, and overall results tend to be disappointing. Studies that have compared acupuncture at theoretically correct sites versus "incorrect" or sham sites have generally found no differences in outcome.

Nonprofit and Proprietary Programs

The oldest of the nonprofit programs is the Five-Day Plan sponsored by the Seventh-Day Adventist Church. ⁸³ An estimated 14 million smokers in more than 150 countries have attended Five-Day Plans. This program considers both

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physical and psychological aspects of cigarette addiction but uses few cognitive-behavioral strategies. Treatment consists of five 90-minute to 2-hour sessions on consecutive days. The Five-Day Plan has been recently revised and renamed the Breathe Free Plan to Stop Smoking, which includes eight sessions during a 3week period.

Both the American Cancer Society and the American Lung Association offer formal group programs. Lando and associates 84 compared these two programs. Smokers (n = 1041) in three Iowa communities were randomly assigned to American Cancer Society clinics, American Lung Association clinics, or an intensive multicomponent behavioral program derived from laboratory research. Although results initially favored the laboratory program over both nonprofit clinics, by 1-year followup, differences between the laboratory program and the American Lung Association program were no longer significant. Sustained abstinence rates at 1 year were 22.2%, 19.0%, and 12.1% for the laboratory, American Lung Association, and American Cancer Society clinics, respectively. The current American Cancer Society Quitline 85 and the American Lung Association smoking cessation programs 86 both offer assistance for those wishing to quit.

A number of commercial programs are available, usually concentrated in larger metropolitan areas. Most programs tend not to be profitable and therefore do not remain active. In evaluating commercial methods, it again appears that the most successful are those that include multicomponent cognitive behavioral techniques. A number of commercial products (eg, lozenges, filters) have been introduced as aids to smoking cessation. Currently, none of these products other than nicotine replacement, bupropion (Zyban), and vareni cline (Chantix) are recognized as effective.

Self-Help

Simply handing smokers written self-help materials has not been demonstrated to be effective. There is evidence based on a limited number of studies that smoker-initiated calls to telephone hotlines or helplines for cessation counseling or assistance do improve abstinence rates. Although results have been mixed, several studies have found good results for proactive telephone support in which calls are initiated by the helpline rather than by the smoker. 87

Computer-Tailored Messages

Computer-tailored messages or "expert systems" have the potential to individualize cessation content to the individual smoker. Some encouraging preliminary results have been reported with these types of programs. 88,89

Pharmacological Intervention

A number of pharmacological aids are recognized as effective by the Food and Drug Administration. Most of these aids involve some form of nicotine replacement: nicotine patch, nicotine polacrilex (nicotine gum), nicotine nasal spray, nico tine inhaler, and nicotine lozenge. Each of these products has specific advantages and disadvantages. All may result in dependence in

some patients.

The patch is easy to use and needs to be applied only once each day. However, it does not allow flexible dosing (eg, once the patch is placed on the skin, the delivered dose is not controlled by the patient), and delivery of nicotine is relatively slow. The gum allows more flexible dosing but is somewhat more difficult to use correctly. Most gum users overdose with this medication. Nicotine nasal spray also has the advantage of flexible dosing, plus it provides faster delivery of nicotine. However, many users are bothered by initial eye and nose irritation, and frequent use is necessary to obtain adequate nicotine levels. The nicotine 341 inhaler allows flexible dosing and at least partially mimics the hand-to-mouth behavior of smoking. The inhaler also has few side effects. A major limitation, however, may be the need to do far more puffing than on a cigarette. For optimal nicotine dosage, many hundreds of puffs may be needed as opposed to perhaps 200 for a regular pack per day smoker. The nico tine lozenge also allows flexible dosing and is easier than nicotine gum for many

The only Food and Drug Administration-approved non- nicotine medications are bupropion hydrochloride (trade name Zyban) and varenicline (trade name Chantix). Bupro pion is available in tablet form. It appears to act on brain chemistry to bring about some of the same effects that nico tine has when people smoke, although its actions are not fully understood. The product is easy to use and can be combined with nicotine replacement. Preliminary evidence suggests that combination of nicotine patch and bupropion may be more effective than either alone. 90 The main ingredient in bupropion has been available for many years for the treatment of depression under the trade name Wellbutrin. However, bupropion works well in smokers with no symptoms of depression. Bupropion is also sold as an antidepressant (Wellbutrin) and has all of the potential side effects of that class of drugs, including suicidality, 20 depression, anxiety, panic attacks, insomnia, and irritability. Monitoring for these symptoms is recommended along with support for cessation. Seizures are a particular problem with doses above 300 mg/day.

Varenicline is also a pill, available through prescription. It works by blocking nicotine receptors in the central nervous system. It should not be used with other quit smoking products. Common side effects are nausea and insomnia, but serious behavioral side effects are also observed. A comparison of varenicline with bupropion and placebo found this agent more effective in smoking cessation at 24 weeks. 91 Var enicline should be stopped if agitation, depression, changes in behavior, or suicidality are observed.

There is evidence that some combinations of medication may be more effective in producing abstinence. A combination of passive dosing (eg, through nicotine patch) and active dosing (eg, ad libitum use, such as with nicotine gum) has been demonstrated to be more effective than a single form of dosing in isolation. 92-94

Clinical Approaches

The many strategies described provide a diverse set of intervention approaches available for clinicians, but there are some proven basic steps summarized as the 5 A's (Box 20-3). These simple actions, aided by intervention strategies, will aid in patients' cessation attempts.

Community and Public Health Approaches

Less than 1% of all smokers have attended formal group or individual treatment programs. Even if half of these smokers achieved permanent abstinence, the overall impact on smoking prevalence would be modest. The need for a



BOX 20-3 Counseling: 5 A's

Ask Systematically identify all tobacco users at every visit. Advise Strongly urge all smokers to quit.

Attempt Identify smokers willing to try to quit. Assist Aid the patient in quitting. Arrange Schedule follow-up contact.



Interventions in health systems

Clinician advice (brief advice significantly increases quitting) Pharmacological

Work site interventions (convenient access to large population of smokers, opportunities to capitalize on social support) Community programs

Overall mixed results

Quit and Win contests widely disseminated

Policy changes

community and systems changes in addition to treatment programs of varying types and intensities, is long apparent. 95,96 These approaches are summarized in Box 20-4. There is growing defined as those who smoked 25 or more cigarettes per day. evidence of the success of these programs. 96 99

Health Care System

Primary care and other clinicians have unique access to the smoking population. At least 70% of smokers see a physician each year. Smokers cite physician advice to quit as an important -20motivator. 67,100 Brief physician advice alone has been associated with a 30% increase in the probability of quitting . 67 Although absolute abstinence rates were modest, universal application of physician or other clinician advice could have a major public health impact. Combining brief advice with offers of behavioral or pharmacological treatment could further increase the likelihood of

reimbursement for smoking cessation services. Curry and colleagues 101 found that use of smoking cessation services varies with the extent of coverage. Full coverage of both behavioral intervention and nicotine replacement therapy led to the highest rates of use of smoking cessation services and to the greatest impact reduce smoking prevalence and cigarette consumption among on the overall prevalence of smoking.

Work Site Interventions

smokers as well as the opportunity to capitalize on social support in this setting. Some very positive results have been reported for work site interventions, although not all studies have been addressed dietary patterns and smoking. Although reductions in tobacco use were in the predicted direction, differences in tobacco significant.

Community Programs

Heart Health Program failed to obtain differences in smoking began in bars and taverns in prevalence between an intervention and a comparison city. The Minnesota Heart Health Program found mixed but primarily negative results for smoking intervention. The only evidence of a significant intervention effect was for women in cross-sectional survey data. 106

Although the overall impact of these community wide trials on smoking prevalence was disappointing, a useful innovation resulting from these programs was the Quit and Win smoking cessation contest. These contests have been successful at the community level in engaging relatively large proportions of the

smoking population. Contests have also engaged large numbers of nonsmokers in support of smokers' quit efforts and have increased community awareness around issues of quitting. Community contests have enrolled as many as 7% of all eligible smokers and have involved many more in reported quit attempts during the contest period. 107 The Quit and Win contest model has been applied in communities in a number of countries around the world and also in national smoking cessation contests in Europe.

A direct successor to these community trials was the COMMIT project, which focused only on smoking. 108,109 In contrast to the earlier studies, COMMIT randomly assigned communities to intervention or control conditions and included sufficient numbers of communities to allow use of the community as the unit of comprehensive public health approach to smoking, which includes analysis. One community within each of 11 matched community pairs (10 in the United States, 1 in Canada) was randomly assigned to intervention. The initial target of COMMIT was heavy smokers, Intervention channels focused on public education through the media and community-wide events, health care providers, work sites and other organizations, and cessation resources. 108

No differences were found between intervention and comparison communities in quit rates among heavy smokers. There was a significant intervention effect on quitting in light to moderate smokers. ¹⁰⁸ Evaluation of overall smoking prevalence failed to indicate significant differences between intervention and comparison communities, however. 109 Results did indicate significant but overall modest differences between smokers in intervention and comparison communities in receipt of intervention activities.

More promising results have been reported for the American A major issue in health system implementation is the lack of Stop Smoking Intervention Study (ASSIST). ASSIST is the largest tobacco control project ever undertaken in the United States. 110 In this initiative, 17 states funded through ASSIST were compared with 32 others (California, which already had extensive tobacco control activities, was omitted). The primary goal of ASSIST was to adults in ASSIST states.

ASSIST was designed as a collaborative effort between the Work sites provide convenient access to a large population of National Cancer Institute and the American Cancer Society and was implemented by state health departments. ASSIST targeted those considered at higher risk for smoking, including youth, ethnic minorities, blue-collar workers, employed, women, heavy smokers, successful. Jeffery and colleagues 102 found modest but significant and smokeless tobacco users. Interventions were delivered to target reductions in overall work site smoking prevalence with an populations through five channels: community environment, work intervention that provided structured group programs and sites, schools, health care settings, and community groups such as incentives (eg, refundable payroll deductions) for quitting. The churches and chambers of commerce. Emphasis was primarily on Working Well Trial conducted in 111 work sites was the largest policy and media interventions, with less emphasis on work site cancer control trial in the United States. 103 Interventions programmatic services. Per capita consumption was almost identical in intervention and comparison states before 1993, when full funding for ASSIST interventions began. By 1996, smokers in use between intervention and control work sites were not the intervention states were consuming approximately 7% fewer cigarettes per capita. 111

Policy Changes

Several major community smoking interventions were offered as Perhaps one of the largest impacts on society may be the part of multicomponent heart disease prevention studies. 49,104,105 improvement in health that results from policy changes, such as the The Stanford Five-City project reported significant smoking passage of smoke-free areas through anti-tobacco legislation or reductions in the intervention cohort relative to the control, but increases in cigarette taxes. An excellent example of this stems from changes were not found in cross-sectional samples. The Pawtucket a study of respiratory health before and after recent prohibition

California. 112 In a small study of 53 bartenders in San Francisco, of 9. those with respiratory symptoms at baseline, 59% no longer had symptoms at follow-up; of those with sensory irritations, 78% had resolution of symptoms (both P < 0.001). Furthermore, a significant improvement (increase) in mean forced vital capacity (increase of 4.2%) was reported after prohibition, and after cessation, significant 11. improvements in both forced vital capacity (6.8% increase) and 12. mean forced expiratory volume in 1 second (4.5% increase) were reported.

Since these early efforts, based on increased cigarette taxes in California, there have been many policy changes built around smoke-free environments, limiting youth access, and tax increases. 15. Pooling Project Research Group: Relationship of blood pressure, serum cholesterol, smoking habit, Environmental changes include smoke-free buildings, transportation, bars, restaurants, and even outdoor places. 97,99,113 ¹¹⁶ Much of the evidence supporting these changes is summarized in an Institute of Medicine report. 117 There is evidence that smoking 17. bans work synergistically with cessation programs. 118 Restricting access of youth to tobacco purchases by enforcing age-related laws is also a successful strategy. 119,120 Finally, increases in tobacco excise taxes have had an effect, especially among younger smokers. 121-124

CONCLUSION

The evidence linking tobacco use to the incidence of and mortality from cardiovascular diseases is substantial. Approximately a halfmillion deaths annually are attributed to cigarette smoking, and the 22. economic costs in medical expenses and indirect costs are enormous. Environmental tobacco smoke is also an important 23. culprit, responsible for some 35,000 to 40,000 deaths from heart disease annually. Important interventions among youth include school-based prevention programs, community-based prevention programs, state and federal initiatives, and cessation assistance. For adults, various behavioral treatments, self-help approaches, and 25. pharmacological therapy are available. Community and public health approaches are invaluable. Physicians and other health care providers need to take greater initiative in reviewing and following up on tobacco use and in informing patients about appropriate 28. Glantz SA, Parmley WW: Passive smoking and heart disease: mechanisms and risk. JAMA community and health care resources for those needing help.

Whereas reductions in adult smoking will have the most immediate benefit in terms of reduced hospitalizations from 30. Otsuka R, Watanabe H, Hirata K, et al: Acute effects of passive smoking on the coronary myocardial infarction and stroke, as well as savings associated with a program reducing smoking prevalence by 1% per year, ¹²⁵ the key ³¹. to making future progress is primary prevention of smoking in children and teenagers. The fact that tobacco is dangerous, lethal, 32. US Department of Health and Human Services: Preventing tobacco use among young people: and disabling when it is used as directed has resulted in the passage of legislation to have the Food and Drug Administration take steps to regulate its sale and use. 126

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Diabetes Mellitus

CHAPTER 21

Diabetes and Cardiovascular Disease

M. Odette Gore, Silvio E. Inzucchi, and Darren K. McGuire

- Diabetes mellitus is a major independent risk factor for cardiovascular disease.
- The worldwide prevalence of diabetes is increasing, driven primarily by the rise in type 2
- The pathophysiology of diabetic cardiovascular disease is multifactorial and incompletely understood.
- Lifestyle intervention is fundamental for the prevention of both type 2 diabetes and its cardiovascular complications.
- Glucose control is important for the management of diabetes, but the most appropriate strategies and glucose targets for cardiovascular disease prevention are still uncertain.
- In addition to lifestyle changes, pharmacologic interventions to treat hypertension and dyslipidemia in diabetes are essential for cardiovascular disease prevention.
- The use of antiplatelet interventions to prevent cardiovascular disease is still controversial but recommended in most patients with diabetes.

The incidence and prevalence of diabetes mellitus are on the rise in the United States and globally, almost entirely due to the growing pandemic of type 2 diabetes. 1 Given that diabetes is a major independent risk factor for cardiovascular disease (CVD), in some clinical contexts considered a coronary disease equivalent, 2 the prevention of diabetes and the management of its associated CVD risk factors are of paramount public health importance. This chapter reviews the epidemiology and preventive strategies for diabetes and its associated CVD complications, with special emphasis on type 2 diabetes, which accounts for more than 90% of diabetes cases worldwide. 3,4

EPIDEMIOLOGY OF DIABETES MELLITUS

Definition

Diabetes mellitus is a group of diseases characterized by insufficient production of insulin or by the failure of the body to respond appropriately to insulin, resulting in hyperglycemia. ³ Vascular complications, the main clinical risk associated with diabetes, are classified as microvascular (diabetic retinopathy, nephropathy, neuropathy) and macrovascular (ischemic heart disease, cerebrovascular disease, peripheral vascular disease).

Diagnostic Criteria

The World Health Organization and the American Diabetes Association (ADA) criteria for the diagnosis of diabetes have evolved during recent decades, summarized in Table 21-1. A diagnosis of diabetes is usually based on tests repeated on at least two different days, unless hyperglycemia is unequivocal or the person is symptomatic. 12,13

Classification

Approximately 90% or more of cases of diabetes mellitus are characterized by relative insulin deficiency with a background of insulin resistance and are classified as type 2 diabetes mellitus. 3,4 The etiology of type 2 diabetes is multifactorial, encompassing genetic, environmental, and behavioral factors, the but exact mechanistic underpinning has not yet been determined; a number of predisposing factors for the development of type 2 diabetes are summarized in Table 21-2.

Type 1 diabetes mellitus results from primary beta-cell loss leading to absolute insulin deficiency, representing less than 10% of cases of diabetes mellitus. 3 The etiology of type 1 diabetes is also multifactorial and poorly understood, although an autoimmune component has been implicated in most cases. Other forms of diabetes not covered in this chapter are gestational diabetes and numerous less common causes (eg, monogenic defects in insulin production or action; diabetes secondary to other pathological conditions of the pancreas, such as pancreatitis or tumors; and drugs or chemicals causing beta-cell toxicity). 3

Prevalence Incidence and of **Diabetes Mellitus**

World

More than 180 million people worldwide were estimated by the World Health Organization to have diabetes mellitus in 2008, 4 increasing from an estimated 135 million in 1995 and projected to rise to 366 million by

*Before this 1979 NDDG publication, there were no unified diagnostic criteria for diabetes. ADA, American Diabetes Association; DCCT, Diabetes Control and Complications Trial; FPG, fasting plasma glucose (fasting is defined as no calorie intake for at least 8 hours); NDDG, US National Diabetes Data Group; NGSP, National Glycohemoglobin Standardization Program; OGTT, standardized oral glucose tolerance test, using a glucose load equivalent to 75 g of anhydrous glucose dissolved in water; PG, plasma glucose; WHO, World Health Organization.

TABLE 21—2 Selected Factors Predisposing Risk for the Development of Type 2 Diabetes Mellitus

Prediabetes, defined as impaired glucose tolerance (2-hr plasma glucose concentration, 140 mg/dL [7.8 mmol/L] to 199 mg/dL [11.0 mmol/L] during an oral glucose tolerance test) or impaired fasting glucose (fasting plasma glucose concentration, 100 mg/dL [5.6 mmol/L] to 125 mg/dL [6.9 mmol/L])

Overweight (BMI > 25 kg/m ²) and at least one other risk factor: First-degree relative with diabetes

One or more features of the metabolic syndrome: HDL-C < 35 mg/dL (0.90 mmol/L), triglycerides > 250 mg/dL (2.82 mmol/L), hypertension (> 140/90 mm Hg or receiving therapy for hypertension)

High-risk population (eg, African American, Latino or Hispanic, Native American, Asian American, Pacific Islander)

Physical inactivity

Age > 45 years

Women who were diagnosed with gestational diabetes mellitus or who delivered a baby weighing > 4 kg (9 pounds)

Other clinical conditions associated with insulin resistance (eg, acanthosis nigricans, polycystic ovarian syndrome)

Obesity (BMI > 30 kg/ ∞)

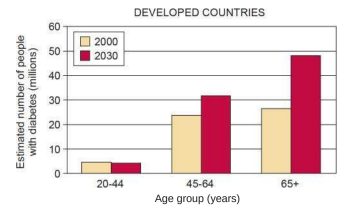
Modified from American Diabetes Association: Standards of medical care in diabetes 2009. Diabetes Care 32:S13, 2009.

2030 (Fig. 21-1). 1 This represents a rise in prevalence adjusted for population growth from 2.8% in 2000 projected to 4.4% in 2030. 1 These numbers probably underestimate the burden of diabetes mellitus in the developing world, where only 25% to 30% of cases are diagnosed. 14

United States

The 2007 diabetes mellitus prevalence estimates for the United States included 17.9 million people with diagnosed

Age group (years)



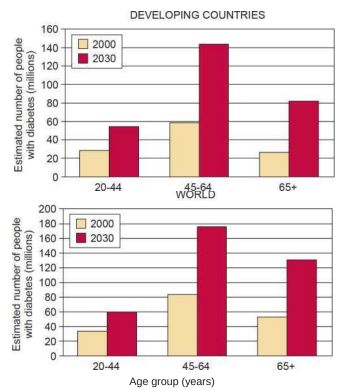


FIGURE 21-1 Estimated number of adults with diabetes by age group, year, and countries for the developed and developing categories and for the world. (From Wild S, Roglic G, Green A, et al: Global prevalence of diabetes: estimates for the year 2000 and projections for 2030. Diabetes Care 27:1047, 2004.)

diabetes and 5.7 million more undiagnosed, representing 7.8% of the US population. ¹⁵ The number of Americans diagnosed with diabetes is projected to increase to 29 million by 2050, with less than one third of this increase attributable to population growth. ¹⁶ More than 1.6 million new cases of diabetes mellitus were diagnosed in the United States in 2007 alone, ¹⁵ representing a marked increase from 625,000 new cases annually in 1990-1992, ¹⁷ even with adjustment for total population (incidence rates 5.3 versus 2.4 per 1000 people per year).

Vulnerable Populations

A number of populations are especially vulnerable to the development of type 2 diabetes. Older persons, women, and especially elderly women are particularly susceptible (Fig. 21-2). ¹ In 2002 in the United States, 1.7% of the population aged 20

FIGURE 21-2 Global diabetes prevalence by sex and age for 2000. (From Wild S, Roglic G, Green A, et al: Global prevalence of diabetes: estimates for the year 2000 and projections for 2030. Diabetes Care 27:1047, 2004.)

to 39 years had diagnosed diabetes compared with 15.1% of those > 60 years. ¹⁸ Diabetes is also more prevalent in certain ethnic groups, including African Americans, Asian Indians, Hispanic or Latino Americans, Native Americans, and Pacific Islanders, among others. For example, the age- and sex-standardized prevalence of diagnosed diabetes reported in NHANES 1999-2002 was almost double in African Americans and Hispanic or Latino Americans compared with non-Hispanic white Americans. ¹⁹ The prevalence of diabetes is high but variable in different Native American populations, ²⁰ reaching 40% to 50% in Pima Indians older than 35 years. ²¹ Somewhat paradoxically, higher socioeconomic status in developing countries and lower socioeconomic status in developed countries have been associated with increased diabetes risk. ^{22,23}

Whereas type 2 diabetes remains less prevalent in children and adolescents than in any other age group, ²⁴ the trend towards increased obesity and decreased physical activity in youth, especially in industrialized countries, is accompanied by an alarming increase in the incidence and prevalence of pediatric type 2 diabetes, ²⁵ especially among ethnic minorities . For example, the population-based multicenter SEARCH for Diabetes in Youth Study reported that among 1530 youth aged 10 to 19 years with newly diagnosed diabetes, type 2 diabetes accounted for 14.9% of diabetes mellitus cases in non-Hispanic white Americans, 46.1% in Hispanics, 57.8% in African Americans, and 86.2% in Native Americans. ²⁴

Factors Contributing to the Increasing Incidence and Prevalence of Type 2 Diabetes Mellitus

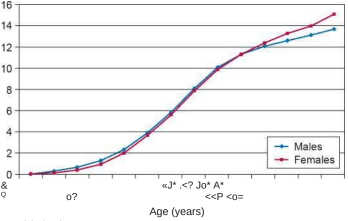
Obesity

The global prevalence of overweight (body mass index [BMI] > 25 kg/m 2 but < 30 kg/m 2) and obesity (BMI > 30 kg/m 2) continues to increase. The World Health Organization estimates that approximately 1.6 billion adults were overweight and at least 400 million were obese worldwide in 2005 and projects an increase to 2.3 billion overweight and 700 million obese by 2015. 26 The risk of diabetes is proportionally increased with both the severity and duration of obesity, as more people are becoming obese earlier in life and rates of extreme obesity (BMI > 35 kg/m 2) has rising. Beyond BMI, measures of increased abdominal obesity, such as waist circumference and waist-to-hip ratio, are even better predictors of diabetes risk.

Diet

It has been difficult to ascertain the role of dietary factors independent of body weight, but the increased availability of

foods high in fat, low in fiber, and with a high glycemic load have been associated with increased risk for type 2 diabetes. ²⁷ Increased consumption of red meats in general and processed meats in particular has also been associated with a higher risk of diabetes. Other dietary factors may also play a role. The rise in diabetes prevalence coincides historically with increasing dietary intake of fructose-rich foods; but in spite of data showing that diets very high in fructose induce insulin resistance and diabetes in laboratory animals, whether high fructose intake plays a role in the pathogenesis of diabetes in humans has not been



established.

Sedentary Lifestyle

In spite of recent indications that leisure-time physical activity may be stable or even increasing in the United States, overall levels have declined significantly during the past half century, and the trend toward an increasingly sedentary life style among Americans continues. 28 Physical inactivity has been associated with an increased risk for developing type 2 diabetes even after adjustment for BMI. Conversely, even modest levels of physical activity intensity and duration are associated with decreased risk. For example, among more than 30,000 women (47% of the Nurses' Health Study cohort) who reported that walking was their only physical activity, diabetes risk decreased significantly across quintiles of energy expenditure, calculated from walking time and pace. ²⁹ In a high-risk population, the Strong Heart Study of 1651 Native Americans reported an odds ratio for incident diabetes reduced by one third (one fourth after adjustment for BMI) in participants who engaged in even modest amounts of physical activity compared with a control group of no physical activity. 30

Aging Population

As the incidence and prevalence of type 2 diabetes increase with age, a global trend towards population aging may account for a portion of the increase in diabetes prevalence. ¹⁶ Life expectancy at birth in developed countries has increased by approximately 25 years in the twentieth century alone, and even though life expectancy in developing countries has increased at a much slower pace, it is projected that about 80% of the world population older than 60 years will live in developing countries by 2050, ³¹ with continued aging of the world population projected through the first half of the twenty-first century.

Diagnostic Criteria and Improved Detection

With the thresholds for diagnosis of diabetes becoming more inclusive in recent years and continued improvements in population-based screening, more persons are being

diagnosed, with relative increases in the ratio of diagnosed to undiagnosed diabetes.

TYPE 1 DIABETES MELLITUS

Definition

Type 1 diabetes results from absolute insulin deficiency due to autoimmune destruction of the insulin-producing beta cells within the islets of the endocrine pancreas. In addition to the obvious glucose abnormalities, patients with type 1 diabetes are also predisposed to ketoacidosis because of unstrained lipolysis and ketogenesis when insulin falls to undetectable or nearly undetectable levels. Indeed, the diagnosis is commonly made in this setting, often in the context of a superimposed intervening illness that increases counterregulatory stress hormones, such as epinephrine and cortisol.

The diagnosis of type 1 diabetes is straightforward in a patient with hyperglycemia (defined according to the criteria in Table 21-1) who is both young and lean, especially one presenting with ketosis or catabolic features, such as weight loss. Serological confirmation with autoantibodies is rarely necessary. Most patients with type 1 diabetes are, however, seropositive for one or more of the following: anti-glutamic acid decarboxylase antibodies, anti-islet cell antibodies 512, and anti-insulin antibodies. When the diagnosis is less clear, such as in an obese child or in a lean older individual, me surement of these serum markers may prove clinically useful.

Type 1 diabetes can be treated only with insulin, and patients are optimally managed with multiple injections per day, involving both basal and prandial components, or continuous subcutaneous insulin infusion (ie, an insulin pump). Oral agents are essentially ineffective and have no standard role in the therapy for this disease.

Epidemiology

The incidence of type 1 diabetes is low, compared with that of type 2 diabetes, and varies widely between populations, 0.1/100,000 per year in China and Venezuela to approximately 37/100,000 per year in Sardinia and Finland.³² Although it is typically diagnosed in children, type 1 diabetes may occur at any age; the rapidity of loss of beta cells is inversely proportional to age, with more gradual development in older individuals during many years. The term *latent autoimmune diabetes of adulthood* has gained favor to describe this form, which initially presents with relatively mild hyperglycemia, often successfully treated with oral agents. Over time, however, insulin deficiency dominates the clinical picture, with labile blood glucose control, insulin dependency, and a propensity for ketosis.

Risk Factors

Although it is clearly of autoimmune origin, the underlying cause or causes of type 1 diabetes remain poorly understood. Defined risk factors include certain HLA types, such as DR3 or DR4, and a family history, although the latter is not as potent a factor as it is in type 2 diabetes. For example, children born to men with type 1 diabetes have about a 6% risk for development of the disease. Children born to women with type 1 diabetes have a lower risk, ranging from 1% to 4%, depending on the age of the mother at delivery. Several viruses have also been implicated in the pathogenesis of type 1 diabetes, including mumps, rubella, cytomegalovirus, and coxsackievirus, but they do not appear to be involved in more than a small minority of cases. The possibilities that dietary factors, micronutrient status, and the individual's intestinal flora may play a role in the predisposition to type 1 diabetes are under active investigation.

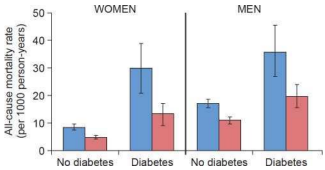
DIABETES MELLITUS AS RISK FACTOR FOR CARDIOVASCULAR DISEASES

Epidemiology

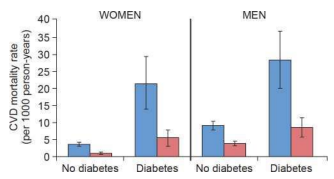
The World Health Organization estimated that as many as 2.9 million deaths worldwide in the year 2005 could be attributed to diabetes, with more than half of these attributable to CVD. ⁴ Of the 284,000 deaths attributable to diabetes in the United States in 2007, about two thirds had CVD as the primary cause of death. ³³ In spite of a trend for reduced all cause and cardiovascular mortality in patients with diabetes in the United States during the past half-century, mortality remains approximately twofold higher in those with versus without diabetes (Fig. 21-3). ³⁴

Diabetic Vascular Disease

Diabetes mellitus is a major risk factor for atherosclerosis, clinically manifested as diabetic macrovascular complications. ³⁵ The risk of ischemic heart disease is twofold to fourfold higher in people with type 2 diabetes compared with those without diabetes, with myocardial infarction (MI) being the number one cause of death. In the Framingham Heart Study between 1976 and 2001, cardiovascular mortality among those with and without diabetes was 6.8 and 2.4 per 1000 person-years, respectively. ³⁴ The risk of MI in patients with diabetes with no history of prior coronary events is, at least in some populations, similar to the risk of MI in patients without diabetes but with prevalent coronary artery disease. ² For example, in a Finnish population-based study of 1059 patients with type 2 diabetes and 1373 without diabetes aged



Note: Bars indicate 95% confidence intervals. Rates are adjusted for age in 10-year



Note: Bars indicate 95% confidence intervals. Rates are adjusted for age in 10-year intervals

FIGURE 21-3 Age-adjusted all-cause (top) and CVD (bottom) mortality rates among participants with and without diabetes by sex and time period. Blue bars represent earlier time period (1950 to 1975); red bars represent later time period (1976 to 2001). (From Preis SR, Hwang SJ, Coady S, et al: Trends in all-cause and cardiovascular disease mortality among women and men with and without diabetes mellitus in the Framingham Heart Study, 1950 to 2005. Circulation 119:1728. 2009.)

45 to 64 years, the 7-year incidence of first MI in those with versus without diabetes was 20.2% versus 3.5%, respectively, and the incidence of recurrent MI was 45% and 18.8%, respectively. 36

This and other similar evidence led the National Choles terol Education Program (NCEP) Expert Panel to recommend that type 2 diabetes be managed as a coronary heart disease equivalent for the purpose of low-density lipoprotein choles -terol (LDL-C) control. ² However, more recent data from clinical trial observations, including studies enrolling patients with newly diagnosed diabetes as well as trials of patients with more advanced type 2 diabetes at high CVD risk, suggest that the CVD risk for diabetes is more intermediate, with risk projected during 10 years ranging between 8% and 20%, ³⁷⁻⁴⁰ contrasted with the "coronary disease equivalent" risk of > 20%/10 years.

Type 1 diabetes is also associated with significantly increased cardiovascular risk, especially in younger patients. Data from the Diabetes UK Cohort, an observational study of 23,751 patients diagnosed with type 1 diabetes in Great Britain, showed that mortality from ischemic heart disease in men and women older than 40 years with type 1 diabetes is increased 4-fold and 7-fold, respectively, compared with the general population; in those younger than 40 years, the risk increase is 9-fold for men and > 40-fold for women. ⁴¹ The Pittsburgh Epidemiology of Diabetes Complications Study, a single-center observational study of 906 patients with type 1 diabetes, ⁴² demonstrated ~15% incidence of CVD after 30 years in this relatively young cohort (median age at onset, 8.5 years).

Having type 2 diabetes doubles one's risk of stroke, even after adjustments for other risk factors (hypertension, dyslipidemia), ⁴³ and increases by 15-fold the risk of lower extremity amputation due to peripheral arterial disease. ⁴⁴ These risks have also been demonstrated in type 1 diabetes, the earlier onset of which results in significantly higher relative risk in younger age groups. For example, the risk of cerebrovascular mortality in the Diabetes UK Cohort was increased by approximately twofold in patients with type 1 diabetes 60 years and older compared with participants without diabetes but by more than fivefold and sevenfold, respectively, in men and women aged 20 to 39 years. ⁴⁵ Finally, atherosclerotic events such as MI and stroke are associated with higher short- and long-term mortality, higher rate of recurrence, and worse overall prognosis in the context of diabetes. ³⁵

Hypertension

Approximately 70% of patients with type 2 diabetes have hypertension (more than double the prevalence in the general population), ⁴⁶ which further increases their CVD risk. For example, an observational analysis of 3642 patients with type 2 diabetes enrolled in the United Kingdom Prospective Diabetes Study (UKPDS) demonstrated a positive adjusted correlation between mean systolic blood pressure and the risk of MI, stroke, and heart failure (Fig. 21-4). ⁴⁷ Hypertension is also an independent risk factor for chronic kidney disease, ⁴⁸ which in turn may exacerbate CVD risk, resulting in a vicious circle. ⁴⁹

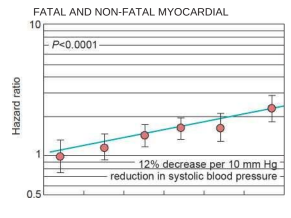
Heart Failure

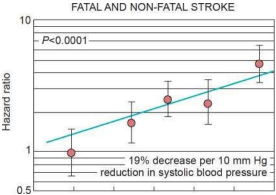
People with type 2 diabetes have a twofold to fivefold increased risk of congestive heart failure (CHF) compared with those without diabetes and have worse outcomes once CHF has developed. ⁵⁰ Diabetes increases the incidence of CHF following the entire spectrum of acute coronary syndromes ⁵¹ and remains an independent predictor of CHF even after adjustment for the increased prevalence of ischemic heart disease among patients with diabetes. ⁵² The increased CHF observed in diabetes is multifactorial and includes more prevalent systolic and diastolic dysfunction due to both

circulatory impairments and derangements of myocardial metabolism (diabetic cardiomyopathy). 53

Sex and Ethnic Differences

Available data suggest that relative cardiovascular risk is higher





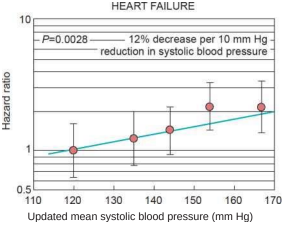


FIGURE 21-4 Hazard rates (95% confidence intervals as floating absolute risks) as estimate of association between category of updated mean systolic blood pressure and myocardial infarction, stroke, and heart failure, with log linear scales. Reference category (hazard ratio 1.0) is systolic blood pressure <120 mm Hg for myocardial infarction and <130 mm Hg for stroke and heart failure; P value reflects contribution of systolic blood pressure to multivariate model. Data adjusted for age at diagnosis of diabetes, ethnic group, smoking status, presence of albuminuria, hemoglobin A1c, high- and low-density lipoprotein cholesterol, and triglyceride. (Modified from Adler AI, Stratton IM, Neil HA, et al: Association of systolic blood pressure with macrovascular and microvascular complications of type 2 diabetes [UKPDS 36]: prospective observational study. BMJ 321:412, 2000.)

in women compared with men with type 1 or type 2 diabetes and that diabetes reduces the sex differences in CVD incidence that otherwise exist in the nondiabetic

Framingham Heart Study increased the risk of CVD mortality by stress such as ischemia, with increased free fatty acid metabolism 3.5- to 5-fold in women and 2- to 3-fold in men. 34

(adjusted hazard ratio [HR], 0.56, 0.68, and 0.68, respectively). metabolic consequences. Asians and Hispanics have a lower risk of stroke and CHF than that of both whites and African Americans (HR, 0.76 and 0.72, stroke; 0.70 and 0.61, CHF). 55 In Pima Indians, despite a very high PREVENTION OF TYPE 2 overall prevalence of diabetes, the incidence of fatal coronary heart **DIABETES MELLITUS** disease is comparatively low. 56

Potential Mechanistic Links Between Diabetes Mellitus and Cardiovascular Disease

Diabetic Macrovascular Disease

The complex mechanistic interrelationships between type 2 diabetes and atherosclerosis have been the focus of extensive Lifestyle Modification and ongoing investigation. A detailed review of this field is beyond the scope of this chapter, but a number of published knowledge in this area. 35,57,58 Among the principal vascular endothelial dysfunction, driven largely by dysregulated nitric interactions, among others. 57.59

Compounding the direct vascular effects of diabetes are a platelet biology yielding a constitutive prothrombotic milieu. 35,57,60 These abnormalities include increased circulating tissue activator inhibitor 1, with decreased levels of antithrombin III placebo control, metformin, or an intensive and protein C. In addition, disturbances of platelet activation, aggregation, morphology, and life span have been well described, further contributing to increased thrombotic potential as well as acceleration of atherosclerosis. 61

Finally, diabetic dyslipidemia is a major contributor to the pathogenesis and progression of atherosclerosis. 62-64 Diabetic a constellation of dyslipidemia is metabolically interconnected lipid and lipoprotein abnormalities, including increased plasma triglycerides, decreased high-density lipo protein (HDL), and a modest increase in low-density lipoproteins (LDL), with larger proportions of atherogenic LDL such as small dense LDL and oxidized LDL. Decreased HDL impairs reverse cholesterol transport (the movement of cholesterol from peripheral tissues to the liver) and reduces the anti-inflammatory and antioxidant effects of HDL in the circulation. 65 Elevated levels of apolipoprotein B-containing lipoproteins, such as LDL and triglyceride-rich very-lowdensity lipoproteins (VLDL), and increased remnant lipoproteins (produced by the hydrolysis of VLDL and chylomicrons) directly promote atherosclerosis. 63

Diabetic Cardiomyopathy

pathogen esis of diabetic cardiomyopathy and CHF. 53 Principal among these processes are abnormal insulin action at the level of the cardiac myocyte coupled with increased circulating free fatty acids, resulting in aberrant myocardial metabolism with accumulation of free fatty acids and triglycerides, and the generation of reactive oxygen species and toxic lipid metabolites, termed cardiac lipotoxicity.

In addition, diabetes is associated with impaired myocellular metabolic substrate switching, deleteriously disturbing the balance

350 population. 34,41,54 For example, type 2 diabetes in the between free fatty acid and glucose metabolism under periods of increasing myocardial oxygen consumption. Hyperglycemia may The risk for diabetic complications is also not uniform exacerbate these effects by inhibiting free fatty acid oxidation, with across ethnic groups. Data from an observational study of 62,432 excess intracellular glucose resulting in nonenzymatic protein diabetic patients showed that African Americans, 21 Asians, and glycation, formation of intracellular and extracellular advanced Hispanics have a relatively lower risk of MI than that of whites glycation end products, and resulting adverse mechanical and

The first line of defense against diabetic CVD is prevention of diabetes. The identification of modifiable risk factors and populations at increased risk for development of type 2 diabetes has made diabetes prevention or delay theoretically feasible. Preventive measures should specifically target people at high risk (see Table 21-2) but could also be applied to the general population. 66

Weight Loss

review articles provide excellent summaries of the current A combined intervention consisting of diet, physical activity, and weight loss reduces diabetes risk, as summarized in Figure 21-5. The disturbances associated with type 2 diabetes is increased Finnish Diabetes Prevention Study included 522 overweight subjects with impaired glucose tolerance ran domized to either oxide biology, indirect and direct vascular effects of advanced usual care or intensive lifestyle interven tion (aimed at reduction of glycation end products, direct adverse effects of increased total intake of fat to < 30% and saturated fat to < 10% of energy circulating nonesterified free fatty acids, and increased consumed, increase in the intake of fiber to > 15 g per 1000 kcal, systemic inflammation and aberrant leukocyte-endothelial moderate exercise for at least 30 minutes per day, and at least 5% weight reduction). 67 After a median follow-up of 4 years, the relative risk of diabetes was reduced by 58% in the intervention number of disturbances in the proteo-fibrinolytic system and group, with sustained benefit observed up to 3 years after the study.

The US Diabetes Prevention Program randomized 3234 people factor, factor VII, von Willebrand factor, and plasminogen with impaired glucose tolerance or impaired fasting glucose to

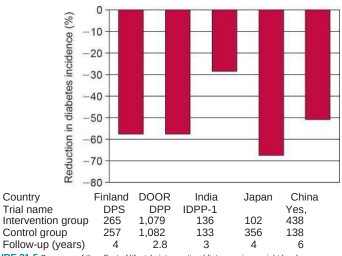


FIGURE 21-5 Summary of the effect of lifestyle intervention (diet, exercise, weight loss) on diabetes incidence in randomized controlled trials (*the Da Qing study was cluster randomized). For Several pathophysiological processes, individually or in - each study, the difference between groups was statistically significant. DPP, US Diabetes Prevention combination, have been proposed to contribute to the Program; DPS, Finnish Diabetes Prevention Study; IDPP-1, Indian Diabetes Prevention Program.

lifestyle modification program aimed at a minimum of 150 minutes versus 6.2%; 37.3% relative risk reduction). 81 of exercise per week, low-fat diet, and 7% weight loss. The incidence of diabetes was reduced by 58% in the lifestyle modification group anatomy through a variety of techniques to reduce calorie intake and by 31% in the metformin group after an average follow-up of and to affect weight loss, has also been shown to have myriad 2.8 years. 69

Program reported a 28.5% reduction in progression to diabetes with reduction, bariatric procedures are associated with beneficial effects lifestyle modification (diet and exercise) for 3 years in Asian Indians on the secretion of nutrient-responsive gut hormones, including with impaired glucose tolerance, 70 and a randomized trial of incretins, ghrelin, and peptide YY, that may improve insulin Japanese men with impaired glucose tolerance observed for 4 years responsiveness independent of weight loss. 82 reported a 67.4% reduction in the risk of diabetes with diet, exercise, and sustained weight loss. 71 The cluster-randomized (by clinic effective strategy for weight loss in severely obese patients. The rather than by participant) Chinese Da Qing study found that at-risk prospective controlled nonrandomized Swedish Obese Subjects individuals in the combined intervention groups (diet, exercise, and trial reported mean weight reductions of 25% for the Roux-en-Y diet plus exercise) had a 51% lower incidence of diabetes compared procedure and 13% to 16% for the restrictive procedures at 10 with the control group during 6 years of active intervention and a years after intervention, compared with a trend for further 43% lower incidence during a 20-year period including 14 years of weight gain in the control group (nonsurgical, nonstandardized post-trial follow-up. 72,73 Despite the variability in risk reduction obesity management). 83 The incidence of diabetes at 2 and 10 across these studies, each has demonstrated that a combination of years was 1% and 7%, respectively, in the overall bariatric diet, physical activity, and weight loss significantly reduces or surgery group, compared with 8% and 24%, respectively, in the delays the development of type 2 diabetes, with several showing control group, with suggestions of modest long-term mortality sustained effects. On the basis of the available evidence, the ADA benefit. 84,85 However, the potential benefits of bariatric surgery recommends reduced intake of dietary calories and fat, regular should be carefully weighed against its significant risk for moderate-intensity physical activity (> 150 min/week), and perioperative complications (including early death) and longsustained weight loss (> 7% of body weight) for diabetes prevention term gastrointestinal and nutritional adverse effects. in people at risk. 74

Independent of weight effects, dietary composition also plays a role in type 2 diabetes risk. Overall, diets high in red and processed meats and saturated fats have been associated with increased risk for type 2 diabetes, and conversely, low-saturated fat, high-fiber Mellitus diets containing whole grains, fruits, vegetables, and fish have been Metformin associated with reduced risk of type 2 diabetes. 67,69,75 The dietary composition recommended by the ADA for diabetes prevention includes reduced intake of fat, increased dietary fiber (14 g fiber per 1000 kcal), and whole grains (at least half of grain intake), and foods with low glycemic index (generally less processed) are encouraged.

Physical Activity and Exercise

Even though physical activity (overall) and exercise (planned and metformin was indistinguishable from placebo in participants structured leisure-time physical activity) may have only a modest older than 60 years as well as in those whose fasting glucose effect on weight if they are not accompanied by dietary measures, 76 concentration was < 110 mg/dL and in those whose BMI was < increasing physical activity, which directly improves insulin 35 kg/m². sensitivity in skeletal muscle, 77 may be beneficial for diabetes prevention independent of weight loss. Prospective cohort studies ("masking" of diabetes) or sustained (true prevention), 1274 of 21,271 American men aged 40 to 84 years participating in the participants who had not developed diabetes by the end of the Physicians' Health Study and of 7735 British men aged 40 to 59 years, Diabetes Prevention Program were subjected to an oral glucose all free of diabetes at baseline, showed that individuals who engaged tolerance test 1 to 2 weeks after the discontinuation of study in moderate levels of physical activity had a significantly lower risk medication. There was an increased incidence of diabetes in the for diabetes compared with those who were physically inactive, metformin group during this washout period (indicating that even after adjustment for age and BMI. 78,79

diabetic women aged 40 to 65 years observed for 8 years) showed with metformin remained significant at 25%. 86 that higher quintiles of physical activity were associated with decreased diabetes risk, and the trend remained significant after drug recommended by the ADA to be considered for diabetes adjustment for BMI. 29

(at least 30 min/day on most days or 150 min/week as tolerance who are younger than 60 years, have a BMI > 35 kg/m recommended by the ADA) is important for diabetes prevention. It 2, and have other diabetes risk factors (HbA1c > 6.0%, should be recommended to individuals at risk in addition to other hypertension, low HDL-C, high triglycerides, family history). 12 weight loss strategies. 74.80

Obesity Treatment

When lifestyle modification alone fails to reduce body weight in obese patients, pharmacological or surgical intervention

may be considered. Orlistat is a gastric and pancreatic lipase 351 inhibitor that reduces fat absorption by ~30%. In a 4-year trial of 3305 obese patients randomized to lifestyle changes I plus either orlistat or placebo, orlistat was associated with a I greater weight loss (5.8 kg versus 3.0 kg) and a lower incidence of diabetes (9.0%

Bariatric surgery, which modifies the gastrointestinal tract beneficial effects on measures of glucose and lipid metabolism, In other high-risk populations, the Indian Diabetes Prevention - including reducing the risk of diabetes. 82 In addition to weight

Several studies support the use of bariatric surgery as an

Drug Therapy for Prevention of Diabetes

The most compelling evidence for pharmacological prevention of diabetes comes from a study of 2155 individuals with impaired glucose tolerance randomized to either metformin or placebo in the Diabetes Prevention Program. After a mean follow-up of 2.8 years, the incidence of diabetes was 4.8% in the metformin group versus 7.8% in the placebo group, representing a 31% relative risk reduction. 69 Notably, however,

To determine whether the effect of metformin was transient masking did occur in those cases), but when the washout data Similarly, data from the Nurses' Health Study (70,102 non- - were included in the overall analysis, the diabetes risk reduction

On the basis of these observations, metformin is the only prevention. It is specifically recommended for high-risk patients On the basis of the available evidence, regular physical activity with both impaired fasting glucose and impaired glucose

352*a* -Glucosidase Inhibitors

2125% relative risk reduction . 87 However, the incidence of diabetes of the diabetes prevention trials were designed or powered to assess significant masking component.

Another drug in this class, voglibose, was studied in a randomized trial of 1780 Japanese subjects at high risk for diabetes and was found to reduce the incidence of diabetes by 40% after a MANAGEMENT OF HYPERGLYCEMIA TO PREVENT short mean follow-up of 48 weeks. 88 This trial was stopped early CARDIOVASCULAR DISEASES because of apparent benefit, which is controversial and could in fact overestimate the effects of the active medication. 89 Of note, agents in this class have significant gastrointestinal side effects that limit Disease Risk: Epidemiology their broad appeal.

Thiazolidinediones

nuclear receptor peroxisome proliferator-activated receptor y (PPAR y) that regulates gene transcription, the thiazolidinediones (TZDs), including rosiglitazone and pioglitazone, target the over longer duration, the achieved insulin sensitization may improve beta-cell preservation, which could plausibly enhance their impaired fasting glucose or impaired glucose tolerance (Fig. 21-6). 91,92 The magnitude of the treatment effects in these trials was large, ranging from 55% to 81% relative risk reduction for diabetes, but 1.43) for each 1% increase in HbA1c. commensurate with the risk reduction achieved by lifestyle intervention described before.

Despite the robust effects observed across the TZD drug class and across trials, however, neither of the two currently available HbA1c were associated with increased mortality, including that due TZDs has a product label indication for this purpose,

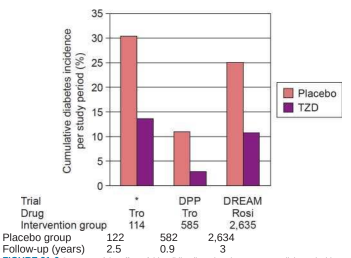


FIGURE 21-6 Summary of the effect of thiazolidinedione (TZD) treatment on diabetes incidence in randomized controlled trials. For each study, the difference between groups was statistically significant. *Hispanic women with previous gestational diabetes; DPP, US Diabetes Prevention Program; DREAM, Diabetes Reduction Assessment with Ramipril and Rosiglitazone Medication; Rosi, rosiglitazone; Tro, troglitazone. 90-92

and they are not recommended for diabetes prevention in society The Study To Prevent Non-Insulin-Dependent Diabetes Mellitus guidelines. 12 The absence of recommendation stems from a number (STOP-NIDDM) randomized 1429 participants with impaired of considerations, including ongoing concerns about adverse effects glucose tolerance to either acarbose or placebo. After a mean follow- such as increased heart failure and fracture risk as well as the current up of 39 months, the cumulative incidence of diabetes was 32% in expense of these drugs. In addition, data are discordant regarding the acarbose group and 42% in the placebo group, representing a the effects of these drugs on atherosclerotic disease risk, 93 and none during 3 months of single-blind post-trial follow-up on placebo effects on macrovascular or microvascular disease endpoints. On the was 45% higher in the former acarbose group, suggesting a basis of these considerations, TZDs are not currently recommended for the prevention of type 2 diabetes.

Glucose and Cardiovascular

Extensive epidemiological evidence has linked measures of hyperglycemia, including fasting plasma glucose, random plasma By virtue of their insulin-sensitizing effects through activation of the glucose, 2-hour post-challenge plasma glucose, and HbA1c, with increased overall and CVD mortality, 94,95 with associations extending well into what is considered the normal range for glucose. One large meta-analysis by Levitan and coworkers 96 of 38 principal pathological underpinning of type 2 diabetes. In addition, investigations found that overall, those with the highest measures of glucose experience approximately one third more CVD events than those with the lowest (RR, 1.36; 95% CI, 1.23-1.52). This relationship effects on prevention or delay of progression to diabetes. 90 Thus, the persisted even when diabetic subjects were excluded from the TZDs appear promising in this regard, with supportive data analysis (RR, 1.26; 95% CI, 1.11-1.43) (Fig. 21-7). Adjustment for CVD deriving from a number of randomized controlled clinical trials risk factors attenuated the relationship to some degree (RR, 1.19; enrolling patients at exaggerated diabetes risk, including Hispanic 95% CI, 1.07-1.32). A second and separate meta-analysis by Selvin women with gestational diabetes, % and large trials of subjects with and colleagues 97 of 13 observational studies of diabetic patients (N = 9123) found a pooled relative risk for CVD of 1.18 in type 2 diabetes (95% CI, 1.10-1.26) and 1.15 in type 1 diabetes (95% CI, 0.92-

> In a more recent analysis using NHANES III data from 19,025 adult participants (baseline survey in 1988-1994 with follow-up through 2000), Saydah and coworkers 98 found that higher levels of to heart disease. After adjustment for other CVD risk factors, the hazard ratio for adults with HbA1c > 8% versus < 6% was 2.59 (95% CI, 1.88-3.56) and 3.38 (95% CI, 1.98-5.77) for all-cause and cardiovascular mortality , respectively. The comparative hazard ratios for adults with a diagnosis of diabetes were 1.68 (1.03-2.74) and 2.48 (1.09-5.64), respectively. However, in this analysis, among those without diagnosed diabetes, no significant association between either all-cause or cardiovascular mortality and HbA1c category was found.

> Within the context of a clinical trial, the largest epidemiological analysis comes from the UKPDS, involving 5102 patients with newly diagnosed type 2 diabetes. In line with the esti mates of Selvin and colleagues, Stratton and collaborators 99 have shown that each 1% reduction in HbA1c was associated with a 14% reduction in MI events. Of note, however, the graded association between HbA1c and microvascular end points was far steeper than that for MI (Fig. 21-8).

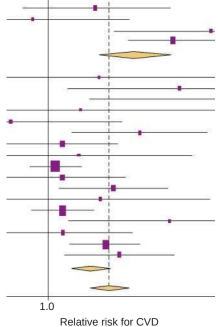
Overview of Current Antihyperglycemic Drug Classes

A variety of both oral and injectable antihyperglycemic agents are used for blood glucose control in patients with type 2 diabetes. In general, these include drugs that stimulate insulin release, improve the body's response to insulin, or delay carbohydrate absorption. In those patients not responding adequately to these agents, typically used in combination,

Cohorts of men

Ohlson et al, 1986 Vaccaro et al, 1992 Yarnell et al. 1994 Haheim et al. 1995 Park et al 1996 Cremer et al., 1997 Folsom et al, men, 1997 Lowe et al, black, 1997 Lowe et al, white, 1997 Balkau et al. 1998 Biornholt et al. 1999 Hart et al, men, 1999 Rodriguez et al, 1999 Wannmethee et al, 1999 Simons et al, men, 2000 DECODE Study group, men, 2001 Henry et al., 2002

Combined



4.0

FIGURE 21-7 Relative risks for CVD comparing the highest with the lowest glycemia categories stratified by sex. Size of the solid squares is inversely proportional to the variance of the study estimate. Arrows represent error bars that continue beyond the scale of the figure, and diamonds represent the random-effects pooled relative risk and 95% confidence interval overall and for analyzes by sex. The dashed line is drawn at the overall pooled estimate. (Modified from Levitan EB, Song Y, Ford ES, et al: Is nondiabetic hyperglycemia a risk factor for cardiovascular disease? A meta-analysis of prospective studies. Arch Intern Med 164:2147, 2004.)

insulin therapy may be used, although patients usually do not require the highly intensive programs needed in those with type 1 diabetes.

05

There are now 11 individual drug categories approved for glucose lowering in patients with type 2 diabetes in the United States. Their mechanisms of action, potency, advantages, and disadvantages are summarized in Table 21-3.

The sulfonylurea drug class is the oldest, in use since the 1950s. Their main side effect is hypoglycemia, which activates an adrenergic response and therefore can be potentially deleterious to the cardiovascular system. Direct negative effects on the heart from these agents have also been proposed because the receptor they activate, the sulfonylurea receptor, is expressed in cardiomyocytes, and activation causes closure of ATPdependent potassium channels (K ATP). Because K ATP closure in myocytes during ischemia may impair preconditioning, sulfonylureas may theoretically exacerbate cardiac injury. Although several animal models have suggested such an effect, the degree to which the drugs bind to K ATP channels in the heart is much less than in pancreatic beta cells, and there are no convincing human data to suggest harm. Sulfo nylureas are therefore widely considered to be safe in diabetic patients, including those with coronary artery disease, although this remains consensus opinion in the absence of rigorous data from clinical outcomes trials. Logically, however, these drugs should not be used in the setting of acute coronary syndromes.

Metformin, a biguanide, reduces glucose primarily by attenuating hepatic glucose output, probably by activation of the enzyme AMP kinase. In the UKPDS randomized trial, the risk of MI was reduced by 39% (P = 0.01) with metformin compared

with those receiving conventional care (diet alone). 100 Such an effect was not convincingly demonstrated in the group assigned to sulfonylureas or insulin (relative risk reduction, 16%; P=0.052). 101 On the basis of these data, paired with a low hypoglycemia risk, modest weight loss, high tolerability with few adverse effects, and low cost, metformin is widely favored as initial monotherapy in most patients with type 2 diabetes. Whereas the use of metformin is cautioned for patients with decompensated heart failure and contraindicated in patients with advanced renal disease because of concern for lactic acidosis, the aggregate data suggest that this risk is quite small or possibly even nonexistent. 102

The TZDs (pioglitazone, rosiglitazone) activate PPAR y and improve insulin sensitivity by increasing glucose uptake by peripheral tissues, mainly skeletal muscle. They also augment lipogenesis by inducing the differentiation of preadipocytes. Their use is associated with a glucose-lowering effectiveness on the same order as that seen with sulfonylureas and metformin and is not associated with hypoglycemia. Pioglitazone, but probably not rosiglitazone, also has beneficial lipid effects, including a 10% to 15% increase in HDL-C and a commensurate decrease in triglycerides. ^{103,104} Unfortunately, TZDs are associated with weight gain and fluid retention as well as increased fracture risk in women. The fluid retention is associated with incident or worsening heart failure. ¹⁰⁵ The incidence of peripheral edema in TZD-treated patients is

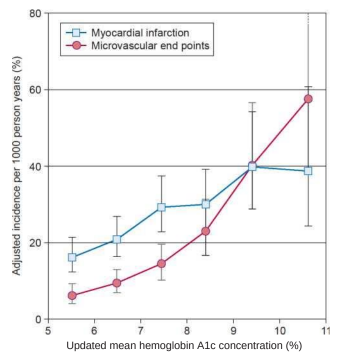


FIGURE 21-8 Incidence rates and 95% confidence intervals for myocardial infarction and microvascular complications by category of updated mean hemoglobin A1c concentration, adjusted for age, sex, and ethnic group, expressed for white men aged 50 to 54 years at diagnosis and with mean duration of diabetes of 10 years. (From Stratton IM, Adler AI, Neil HA, et al: Association of glycaemia with macrovascular and microvascular complications of type 2 diabetes [UKPDS 35]: prospective observational study. BMJ 321:405, 2000.)

approximately 5% to 10%, more so when they are used in conjunction with insulin. The risk of heart failure is approximately 1%, also higher in insulin-treated patients. The relative risk of heart failure in patients taking TZDs is approximately 1.7 compared with a non-TZD regimen, with a higher reported risk increase with rosiglitazone than with pioglitazone. 106,107

There is little cardiovascular information available for the other antihyperglycemic therapies currently marketed for diabetes. There are very few data with the newer incretin based therapies, such as the glucagon-like peptide 1 (GLP-1) agonists and the dipeptidyl peptidase 4 inhibitors. There is great interest in the potential cardiovascular benefit of GLP-1 agonists (eg, exenatide) because these agents result in substantial weight loss in some patients, with the early demonstration of associated improvement in various cardiovascular risk factors. Studies suggesting a benefit on ventricular function in heart failure and after acute coronary events are very preliminary. ¹⁰⁸

Of note, regulatory agencies around the globe have recently begun requiring clinical trial assessment of the CVD effects of diabetes drugs under development, with a requirement to demonstrate a nominal degree of cardiovascular safety before drug approval. ^{109,110}

Type 2 Diabetes Mellitus

UKPDS

The UKPDS demonstrated microvascular risk reduction from more intensive glycemic control (relative risk reduction, 21% to 34%) in 5102 patients with newly diagnosed type 2 diabetes . 101 patients were randomized to treatment with sulfonyl urea or insulin as first-line therapy compared with diet alone. During a mean follow-up of 10 years, mean HbA1c was 7.0% in actively treated patients versus 7.9% in the control group, but the effect on macrovascular outcomes (nonfatal and fatal MI and sudden

death) did not reach statistical significance (16% risk reduction; P = 0.052). In a randomized substudy (N = 753) of the UKPDS, initial monotherapy in overweight patients (> 120% of ideal body weight) with metformin resulted in improved macrovascular outcomes, an outcome not observed within the main trial (see earlier).

From the UKPDS came the original notion that glucose control, although apparently important for microangiopathy, had a more obscure effect on CVD complications. Given the metformin findings, it also raised concerns about whether the method by which glucose is lowered may play some role in altering this disease's cardiovascular risk equation.

Long-term post-trial follow-up of the UKPDS cohort has demonstrated cardiovascular benefit from previous intensive glucose control. Despite a loss in the HbA1c differences between the groups, during 10 years of post-trial monitoring, the patients previously intensively treated with sulfonylurea or insulin experienced relative risk reductions for any diabetes -related endpoint (9%; P = 0.04) and microvascular disease (24%; P =0.001), whereas decreases in the relative risk for MI (15%; P =0.01) and all-cause mortality (21%; P = 0.01) eventually emerged. ³⁹ In the overweight metformin-treated patients, corresponding reductions in any diabetes-related endpoint (21%; P = 0.01), MI (33%; P = 0.005), and all-cause mortality (27%; P = 0.002) were observed. These important results demonstrate a "glycemic memory" concept whereby the effects of intensive treatment continued beyond the ran domization phase of the study, eventually resulting in beneficial macrovascular outcome differences.

ACCORD, ADVANCE, VADT

More recently, three randomized clinical trials (ACCORD, 37 ADVANCE, 40 and VADT 38) formally tested the cardiovascular effect of more versus less intensive glucose control (HbA1c target of 6% to 6.5% versus 7% to 9.0%) during a period of 3 to 5 years, all three failing to demonstrate significant benefit (Table 21-4). In the ACCORD trial, cardiovascular mortality was actually increased in the intensive group, although the explanation for this phenomenon remains enigmatic. Hypoglycemia was more common in patients who died in both intensive and standard control groups, but post hoc analyzes have not been able to demonstrate a causative relationship. Notably, more than three of four intensively treated patients received insulin during the course of the trial, and the majority of patients were prescribed at least three oral agents simultaneously. Whether such polypharmacy may have played a role in the study results is also not clear.

In post hoc analyzes from both ACCORD and ADVANCE, the subgroup of patients without preexisting vascular complications actually showed a benefit from more intensive management, experiencing fewer cardiovascular events. In ADVANCE, the primary endpoint, which was a composite of both microvascular and macrovascular events, was modestly reduced in the intensive arm, although this was driven solely by improved intermediate markers of renal disease (relative risk reduction, 21%; P = 0.006). In a post hoc analysis from the VADT, intensive glycemic control appeared to confer a CVD benefit in patients with duration of diabetes < 12 years but had a neutral or even adverse effect in patients with longer duration of the disease. Post hoc exploratory data from the VADT also suggested that an episode of severe hypoglycemia within 90 days was a strong predictor of CVD events and CVD mortality. Importantly, none of these trials addressed the question of whether more intensive glycemic management may have macrovascular benefits if it is applied and maintained for longer than 5 years or if it is initiated at the time of diagnosis before vascular complications have become established. An even more recent post hoc analysis suggests

TABLE 21-3	Pharmacologic Therap	by for Type 2	Diabetes Mellitus			
Drug Class Sulfonylureas	Examples Glyburide Glipizide Glimepiride	Underlying Mechanism Closes K ATP channels	Main Metabolic Effects T Pancreatic insulin secretion	and A1c ~ 1%-2%	Advantages Microvascular risk	Disadvantages Hypoglycemia Weight gain Ischemic preconditioning (?) Beta-cell exhaustion
Glinides	Repaglinide Nateglinide	Closes K ATP channels	T Pancreatic insulin secretion	~ 1%-1.5%	More physiological than sulfonylureas Postprandial glucose	Hypoglycemia Weight gain Ischemic preconditioning (?) Beta-cell exhaustion Dosing frequency
Biguanides	Metformin	Activates AMP kinase	e Hepatic glucose production	~ 1%-2%	No hypoglycemia Weight loss and CVD events	Gastrointestinal side effects (diarrhea) Lactic acidosis Multiple contraindications to consider
Thiazolidinediones	Rosiglitazone Pioglitazone	Activates PPARy	T Peripheral insulin sensitivity	~ 1%-1.5%	No hypoglycemia Beta- cell preservation and CVD events (?) (pio) HDL -C Triglycerides Blood pressure	Weight gain Edema, heart failure Bone fractures (women) T LDL-C (T particle size) Rosiglitazone controversy in coronary heart disease
α- Glucosidase Inhibitors	Acarbose Miglitol	Small blocks intestinal a - glucosidase	Intestinal carbohydrate absorption	~ 0.5%-1%	No hypoglycemia Non-systemic Postprandial glucose and CVD events (?)	Gastrointestinal side effects (flatulence) Dosing frequency
Glucagon-like peptide 1 (GLP-1) agonists	Exenatide	Activates GLP-1 receptors	T Pancreatic insulin secretion i Pancreatic glucagon secretion Delays gastric emptying T Satiety	~ 1%	Weight loss No hypoglycemia Betacell preservation Postprandial glucose Cardiovascular benefits (?)	Gastrointestinal side effects (nausea, vomiting) Pancreatitis (?) Injectable
Amylinomimetics	Pramlintide	Activates amylin receptors	i Pancreatic glucagon secretion Delays gastric emptying T Satiety	~ 0.5%	Weight loss Postprandial glucose	Gastrointestinal side effects (nausea, vomiting) Dosing frequency Injectable
Dipeptidyl peptidase (DPP) 4 inhibitors	Sitagliptin Vildagliptir Saxagliptin	Inhibits DPP-4 T Endogenous incretin levels	T Pancreatic insulin secretion i Pancreatic glucagon secretion	~ 0.6%-0.8%	No hypoglycemia	Urticaria, angioedema, pancreatitis (?)
Acid balls sequestrators	Colesevelam	Binds bile acid cholesterol	?	~ 0.5%	No hypoglycemia and LDL-C	Gastrointestinal side effects (constipation) T Triglycerides Dosing frequency
Dopaminergic receptor 2 (D ₂) agonists	Bromocriptine	Activated dopaminergic receptors	Modulates hypothalamic circadian organization Hepatic glucose production	~ 0.5%	No hypoglycemia	Gastrointestinal side effects (nausea) Dizziness
Insulin	Human NPH Regular human Glargine, detemir Lispro aspartame Glulisine Premixed (various types)	Activates insulin receptors	T Peripheral glucose disposal Hepatic glucose production and Proteolysis Lipolysis Ketogenesis	No limit	Microvascular risk Universal effectiveness	Hypoglycemia Weight gain Injectable Training requirements "Stigma"

TABLE 21—4 Baseline Characteristics and Main Results from Three Large Randomized Cardiovascular Trials in Patients with Type 2 Diabetes Mellitus

	AGREEMENT 37		ADVANCE 40		VADT ³⁸	
N	10,251		11,140		1791	
Age (mean, years)	62		66		60	
BMI (mean, kg/ m²)	32		28		31	
Follow-up (mean, years)	3.5		5		5.6	
HbA1c target	< 6.0% vs. 7.0%-7	.9%	< 6.5% vs. "stand	ard"	< 6% vs. 8%-9%	
Baseline HbA1c (mean)	8.3%		7.5%		9.4%	
Endpoint HbA1c (mean)	Intensive 6.4%	Standard 7.5%	Intensive 6.43%	Standard 7.0%	Intensive 6.9%	Standard 8.4%
Severe hypoglycemic events	Intensive 10.5%	Standard 3.5%	Intensive 2.7%	Standard 1.5%	Intensive 8.5%	Standard 2.1%
Weight change	Intensive + 3.5 kg	Standard + 0.4 kg	Intensive - 0.1 kg	Standard - 1.0 kg	Intensive + 8.1%	Standard + 4.1%
Major macrovascular or microvascular event	Not reported		0.9 (0.82-0.98);	P = 0.01	0.88 (0.74-1.0	5); <i>P</i> = 0.14
Nonfatal MI or stroke, cardiovascular death	HR, 0.9 (0.78-1.04); P = 0.16	0.94 (0.84-1.06);	; P = 0.32	Not reported	
All-cause mortality	HR, 1.22 (1.01-	-1.46); <i>P</i> = 0.04	0.93 (0.83-1.06);	; P = 0.28	1.07 (0.81-1.4)	2); <i>P</i> = 0.62
Nonfatal MI	HR, 0.76 (0.62-0.9	2); <i>P</i> = 0.004	0.98 (0.77-1.22);	; P = NS	0.82 (0.59-1.14	4); <i>P</i> = 0.24

unsuccessful intensive glycemic control (the failure to respond to the intervention) was linked to the increased car diovascular disease mortality seen in the intensive group. ^{110a}

A rational synthesis of the data has led the ADA to maintain its general treatment target of HbA1c < 7%, primarily to minimize microvascular disease risk. ^{12,111} The data in support of this target for macrovascular risk reduction are uneven at best, and there is little evidence to support any lower target from a CVD standpoint. In fact, the ADA statements have considered that a less stringent target may be reasonable for diabetic patients with a history of severe hypoglycemia, advanced complications, or comorbid conditions or for those who are difficult to control or who have had diabetes for many years. Importantly, as described later, meticulous attention to other cardiovascular risk factors, such as lipids and blood pressure, has a much more prominent role than glucose lowering in this realm.

Thiazolidinedione Studies: PROactive, RECORD

Because of the well-recognized association between insulin resistance and CVD, the development of an insulin-sensitizing class of medications was initially met with great hope that these agents could not only lower glucose but also directly reduce CVD complications. Indeed, preliminary in vitro and animal investigations suggested a potent antiatherosclerotic effect of the TZDs, results that were subsequently confirmed in humans by use of surrogate endpoints, such as coronary and carotid atherosclerosis as measured by ultrasound as well as intracoronary stent restenosis. 112,113 However, it has been more challenging to confirm an actual benefit on cardiovascular events.

In the Prospective Evaluation of Pioglitazone and Macrovascular Events (PROactive), 5238 patients with type 2 - diabetes and established macrovascular disease were assigned to pioglitazone versus placebo, added to their prior antihyperglycemic regimen. ¹⁰³ The primary endpoint, a broad CVD composite, occurred in similar numbers of those assigned to active therapy versus placebo (21% versus 23.5%; HR, 0.90;

95% CI, 0.80-1.02). In the analysis of the main secondary composite endpoint of mortality, MI, and stroke, however, a significant benefit was apparent in patients receiving pioglitazone (12.3% versus 14.3%; HR, 0.84; 95% CI, 0.72-0.98) (Fig. 21-9). This apparent benefit was to some degree attenuated by an increased risk of heart failure hospitalizations in active therapy patients (5.7% versus 4.1%), although no measurable increase in heart failure mortality was observed. 105 Pioglitazone compared with placebo was also associated with improvements in several CVD risk factors, including HbA1c, blood pressure, HDL-C, and triglycerides, that could individually or aggregately contribute to the observed effects on atherosclerotic disease endpoints. A direct effect of pioglitazone on atherosclerosis is further supported by results from two randomized activecontrolled trials using ultrasound assessments of carotid and coronary arteries; favorable effects of pioglitazone compared with glimepiride on these imaging intermediates were demonstrated in both trials. 112,113 In spite of these results, it is not yet possible to ascribe the proposed antiatherosclerotic effect of pioglitazone to a *direct* effect at the level of the vasculature or, for that matter, to a reduction in insulin resistance. Notwithstanding, in retrospect, the positive findings from PROactive stand alone to some degree, in light of the ACCORD, ADVANCE, and VADT results. They also indicate that the theory that improving insulin sensitivity may decrease cardiovascular events remains worthy of further study.

The other TZD, rosiglitazone, has had an even more controversial history. In 2007, a widely publicized meta-analysis suggested that rosiglitazone was associated with an *increase* in MI risk (HR, 1.43; 95% CI, 1.03-1.98) and a trend toward increased cardiovascular mortality (HR, 1.64; 95% CI, 0.98 2.74). ¹¹⁴ Subsequently, in the RECORD trial involving 4447 patients with type 2 diabetes in whom rosiglitazone was added to either metformin or a sulfonylurea, compared with patients receiving both traditional agents, the hazard ratio for the primary composite CVD endpoint was 0.99 (0.85-1.16). ¹¹⁵ On the basis of these data, the effect of rosiglitazone on ischemic heart disease

risk remains uncertain. In 2010, given the uncertain CVD safety of rosiglitazone, US and European $\,$

FIGURE 21-9 Kaplan-Meier curve of time to the main secondary endpoint, death from any cause, nonfatal myocardial infarction (excluding silent myocardial infarction), or stroke, from the PROactive randomized trial of pioglitazone versus placebo in 5238 patients with diabetes and cardiovascular disease. (From Dormandy JA, Charbonnel B, Eckland DJ, et al: Secondary prevention of macrovascular events in patients with type 2 diabetes in the PROactive Study [PROspective pioglitAzone Clinical Trial In macroVascular Events]: a randomized controlled trial. Lancet 366:1279, 2005.)

regulatory agencies acted to suspend rosiglitazone Number of risk from the European market and severely restrict its use Pioglitazone in the United States. The disparate CVD outcomes with rosiglitazone and pioglitazone could in part be explained by the differential effects these two TZDs have on lipid metabolism.

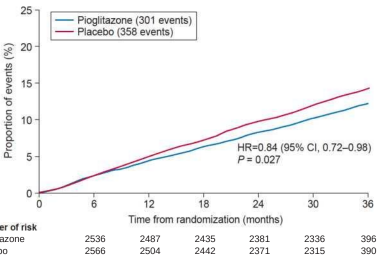
The BARI 2D study demonstrated no benefit on CVD events in a group of 2368 coronary artery disease patients with type 2 diabetes from a glucose-lowering strategy that emphasized insulin sensitization (metformin or TZD) versus one that focused on insulin provision (sulfonylurea or insulin). 116 However, the results of this study cannot be considered conclusive because the TZD in this study was rosiglitazone, and there was a fair degree of crossover therapy in both groups, especially in those randomized to the insulin sensitizers. BARI 2D also explored prompt versus deferred revascularization in a 2 x 2 factorial design. From post hoc analyses, in those patients assigned to prompt revascularization, the insulin sensitizer strategy was associated with a trend toward a lower rate of major cardiovascular events (20.3% versus 25.2%; P = 0.059). This appeared to be driven mainly by a strong trend in the strata undergoing coronary artery bypass grafting (18.7% versus 26%; P = 0.066).

Societal Guidelines for Antihyperglycemic Therapy

As noted before, the ADA continues to recommend that patients with diabetes have their HbA1c controlled to < 7%. This professional organization does not advise a specific method to lower glucose, appreciating the fact that individualization is important. Specifically, older patients with overt CVD did not benefit from the more stringent glycemic policies in ACCORD, ADVANCE, and VADT and may be at increased risk from hypoglycemia or polypharmacy. Accordingly, less rigid targets may be appropriate for this group. A consensus statement sponsored by the ADA and the European Association for the Study of Diabetes was published in 2004 and revised in 2007 and again in 2009. 117 The current cardinal recommendations from this group include the initiation of metformin as foundation therapy in all patients, proceeding expeditiously to combination therapy once glycemic goals are no longer being attained, including consideration of the early use of insulin. The proposed algorithm is shown in simplified form in Figure 21-10.

These recommendations are not considered to be official positions by the organizations, and there remains a significant variability in practice, especially regarding which drug to add after metformin monotherapy. The American Association of Clinical Endocrinologists has proposed a "road map" for the treatment of type 2 diabetes. This is, generally speaking, more inclusive of some of the newer therapies (Fig. 21-11). 118

In all, the simplest regimen that is effective in lowering glucose, that is acceptable to the patient, and that minimizes side effects, especially hypoglycemia, is likely to be the best program for an individual patient. From a CVD standpoint, the early use of metformin is most justifiable. The precise form of



antihyperglycemic therapy beyond metformin remains arguable. The central points to consider are the risk of heart failure with TZDs, the possible CVD risk with rosiglitazone and CVD benefits of pioglitazone, the risk of hypoglycemia with both insulin and insulin secretagogues, and the potential emerging benefit in the form of weight reduction from the newer GLP-1-based therapies.

Vascular Complications of Type 1 Diabetes Mellitus and Their Relationship to Glucose Control

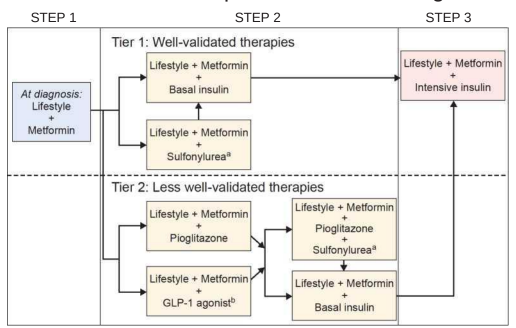
Type 1 diabetes mellitus is associated with both microvascular and macrovascular complications, with a clear relation to the quality of glucose control achieved. The Diabetes Control and Complications Trial (DCCT) assessed the effect of tighter versus conventional glucose management with insulin therapy in 1441 patients with type 1 diabetes, mean age ~27 years, for a mean of 6.5 years. 119 It compared the more stringent strategy of three or four insulin injections per day or an insulin pump and monitoring of blood glucose concentration at least four times daily with conventional treatment of one or two injections per day and just daily monitoring. The glycemic goals in the intensive group included a preprandial blood glucose concentration of 70 to 120 mg/dL, postprandial blood glucose concentration of < 180 mg/dL, and HbA1c level of < 6.05%. In the conventional group, there were no specific glycemic targets, the primary goal being to avoid symptomatic hyperglycemia and hypoglycemia. The mean HbA1c level at the end of the trial in those randomized to intensive therapy was 7.4% compared with 9.1% in the conventional therapy group.

The relative risk for the development or progression of microvascular complications (retinopathy, nephropathy, or neuropathy) was reduced by 54% to 63% in the more tightly managed cohort (Figs. 21-12 to 21-14). Cardiovascular events, however, were few in these relatively young patients. Whereas there were less cardiovascular and peripheral vascular events in intensively treated individuals, the results did not achieve statistical significance (0.5 versus 0.8 events per 100 patientyears; relative risk reduction, 41%; 95% CI, - 10% to 65%).

At the conclusion of the trial, the DCCT patients continued to be observed in the nonrandomized, uncontrolled



2009 ADA / EASD Proposed Consensus Algorithm

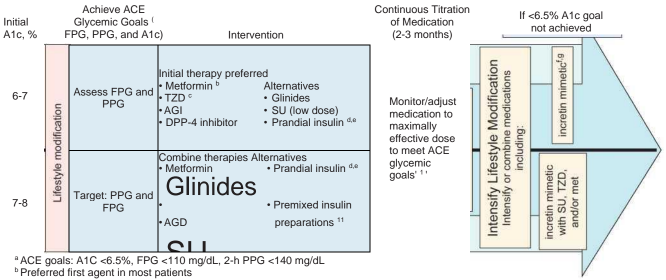


Reinforce lifestyle changes at every visit and check A1C every 3 months until < 7.0%, then at least every 6 months thereafter. Change interventions whenever A1C > 7.0%.

^a Sulfonylureas other than glibenclamide (glyburide) or chlorpropamide. ^b Insufficient clinical use to be confident regarding safety.

FIGURE 21-10 Algorithm for the metabolic management of type 2 diabetes. The tier 1 algorithm represents the best established and most effective and cost-effective therapeutic strategy for achieving the target glycemic goals. Step 1, Lifestyle interventions should be initiated as the first step in treating new-onset type 2 diabetes, and should be reinforced at every visit. Metformin is recommended as the initial pharmacological therapy, in the absence of specific contraindications, for its effect on glycemia, absence of weight gain or hypoglycemia, generally low level of side effects, high level of acceptance, and relatively low cost. Step 2, If lifestyle intervention and the maximal tolerated dose of metformin fail to achieve or sustain the glycemic goals (or if metformin is contraindicated or not tolerated), another medication should be added within 2-3 months of the initiation of therapy or at any time when the target A1c level is not achieved. The second medication recommended is either insulin or a sulfonylurea, with insulin usually reserved for patients with an A1c > 8.5% or with symptoms secondary to hyperglycemia. Step 3, If lifestyle, metformin, and sulfonylurea or basal insulin do not result in achievement of target glycemia, the next recommended step is initiation or intensification of insulin therapy (with discontinuation of insulin secretagogues). The tier 2 algorithm may be considered in selected clinical settings. Specifically, when hypoglycemia is particularly undesirable (eg, in patients who have hazardous jobs), the addition of exenatide or pioglitazone may be considered. Rosiglitazone is not recommended. If promotion of weight loss is a major consideration and the A1c level is close to target (< 8.0%), exenatide is an option. If these interventions are not effective in achieving target A1c, or are not tolerated, addition of a sulfonylurea could be considered. Alternatively, the tier 2 interventions should be stopped and basal insulin started. (Modified from Nathan DM, Buse JB, Davidson MB, et al: Medical management of hyperglycemia in type 2 diabetes: a consensus algorithm for the initiation and adjustment of therapy: a consensus statement of the American Diabetes Association and the European Association for the Study of Diabetes. Diabetologia 52:17, 2009.)

AACE Road Map for Glycemic Control in Treatment-Naive T2DM Patients



^c According to the FDA, rosiglitazone is not recommended with insulin

FIGURE 21-11 The American Association of Clinical Endocrinologists has proposed this road map for the pharmacological management of hyperglycemia in patients with type 2 diabetes. ¹¹⁸ It is generally more inclusive than other guidelines and is unique insofar as it advises a different approach based on the initial HbA1c level

Epidemiology of Diabetes and Its Complications (EDIC) study. ¹²⁰ All patients were offered intensive management, although the meticulous follow-up by the study investigators and coordinators could no longer be provided. Not surprisingly, within 1 year, the HbA1c distinction between the two groups had disappeared. After 11 years, the mean HbA1c level in the original intensive group had risen to 7.9%, whereas it had fallen in the original conventional cohort to 7.8%.

Importantly, despite what appeared to be similar glucose control for most of the follow-up period, differences in microvascular complications persisted, and to some degree, cardiovascular complications appeared to continue to diverge. Ultimately, there were 46 total CVD events in 31 patients in the intensive group and 98 events in 52 patients in the conventional group, for a relative risk reduction of 42% (95% CI, 9%-63%; P = 0.02) (Fig. 21-15). ¹²¹ The composite risk of car diovascular mortality, nonfatal myocardial infarction, or stroke was decreased by 57% (95% CI, 12%-7%; P = 0.02). Notably, other cardiovascular risk factors, including blood pressure, lipids, and BMI, were similar between the groups. One exception was the degree of albuminuria, which was persistently elevated in the conventional cohort (116 versus 54 mg/24 hr) and proved to be an independent predictor of cardiovascular risk.

This persistence of benefit for microvascular complications and the emergence of apparent benefit for macrovascular complications has been termed the legacy effect. Although it is not yet fully explained biologically, this term suggests a vascular benefit from any sustained period of good glycemic control in previous years. It is not yet known whether patients with type 1 diabetes may incur further benefit from even tighter glycemic control, such as the near-normalization of blood glucose concentration. In light of recent trial results in type 2 diabetes

(see earlier), this would seem unlikely, however, especially because patients with type 1 diabetes are even more prone to hypoglycemia and its sequelae.

Global CVD risk prevention strategies beyond glucose control in patients with type 1 diabetes mirror those interventions used in type 2 diabetes, as discussed next, although there have been no large, randomized clinical trials addressing other specific comorbidities in this group of patients.

GLOBAL MANAGEMENT OF TYPE 2 DIABETES MELLITUS FOR THE PREVENTION OF CARDIOVASCULAR DISEASE

Therapeutic Lifestyle Intervention

Lifestyle intervention is the cornerstone of CVD prevention in type 2 diabetes. The general principles of lifestyle intervention summarized earlier in this chapter for type 2 diabetes prevention are also applicable for the management of diagnosed type 2 diabetes. The goals of lifestyle intervention to prevent cardiovascular complications in patients with type 2 diabetes are improved glycemic control, reduced dyslipidemia and hypertension, and smoking cessation.

Weight Loss

Sustained moderate weight loss is recommended by the ADA in all patients with diabetes who are overweight or obese. ¹² Moderate weight loss has been shown to reduce insulin resistance , dyslipidemia, and blood pressure. ^{2,67,69-71,73,74} Although overweight and obesity were associated with lower mortality

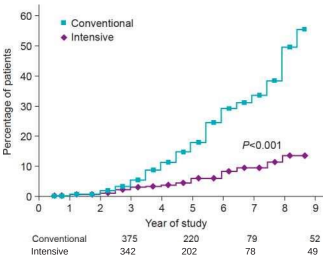
^d Analog preparations preferred

e Rapid-acting analog insulin (available as lispro, aspart, and glulisine) or regular insulin

fundicated for patients not at goal despite SU and/or metformin or TZD; incretin mimetic is not indicated with insulin

⁹ Available as exenatide

^h Available as glargine and detemir



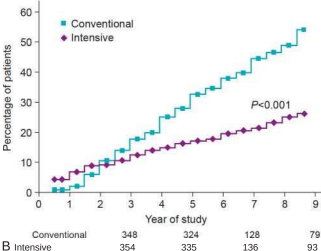
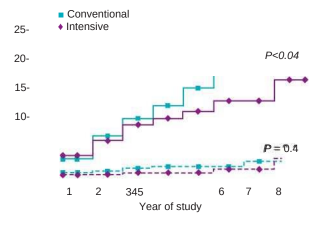


FIGURE 21-12 Microvascular complications in the Diabetes Complications and Control Trial, intensive versus conventional groups. In the primary prevention **(A)** and secondary intervention **(B)** cohorts, intensive therapy reduced the adjusted mean risk of the onset of retinopathy by 76% (P < 0.001) and 54% (P < 0.001), respectively, compared with conventional therapy. The numbers of patients in each therapy group who were evaluated at years 3, 5, 7, and 9 are shown below the graphs. (*From The Diabetes Control and Complications Trial Research Group: The effect of intensive treatment of diabetes on the development and progression of long-term complications in insulindependent diabetes mellitus. N Engl J Med 329:977, 1993.)*



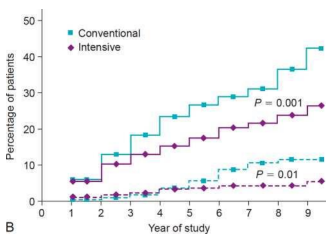


FIGURE 21-13 Microvascular complications in the Diabetes Complications and Control Trial, intensive versus conventional groups. In the primary prevention cohort (**A**), intensive therapy reduced the adjusted mean risk of microalbuminuria by 34% (P < 0.04), (solid lines). In the secondary intervention cohort (**B**), intensive therapy reduced the risks of macroalbuminuria by 56% (P = 0.01) (dashed lines) and microalbuminuria by 43% (P = 0.001) (solid lines). (From The Diabetes Control and Complications Trial Research Group: The effect of intensive treatment of diabetes on the development and progression of long-term complications in insulin-dependent diabetes mellitus. N Engl J Med 329:977, 1993.)

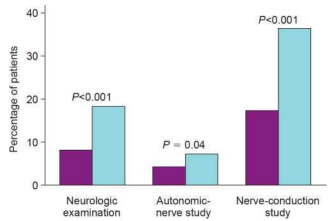


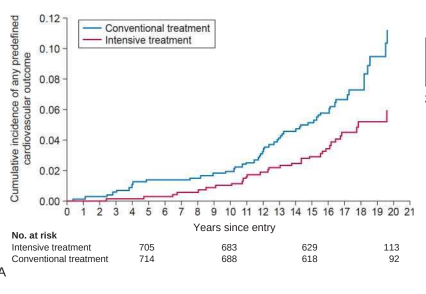
FIGURE 21-14 Microvascular complications in the Diabetes Complications and Control Trial, intensive versus conventional groups. Prevalence of abnormal clinical neurologic examination, autonomic nerve studies, and nerve conduction studies at 5 years between intensive (purple bars) and conventional (blue bars) treatment groups. (From The Diabetes Control and Complications Trial Research Group: The effect of intensive treatment of diabetes on the development and progression of long-term complications in insulin-dependent diabetes mellitus. N Engl J Med 329:977, 1993.)

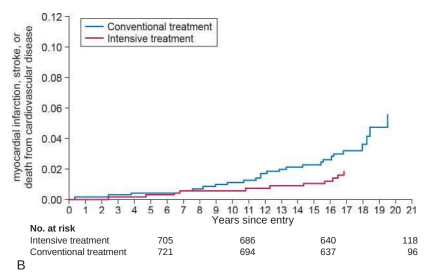
(the so-called obesity paradox) in some studies of patients with established CVD, current evidence supports the role of moderate weight loss in CVD prevention in overweight or obese individuals. 122

The ongoing Look AHEAD (Action for Health in Diabetes) multicenter prospective trial has randomized 5145 overweight or obese patients with type 2 diabetes (mean BMI, 36.0 kg/m 2) to intensive lifestyle intervention (diet and exercise) or control (support and education), aiming to examine the long-term effects of lifestyle intervention on the incidence of major CVD events. 123 The study is scheduled to conclude in 2012, but the 1-year interim results have already shown that intensive lifestyle intervention led to an average 8.6% weight reduction, compared with 0.7% in the control group, and was associated with significant improvement in glycemic control and cardiovascular risk factors (Table 21-5). 123

Continued intervention and follow-up will determine whether these effects are sustained in the long term and are associated with reduced incidence of CVD events. Ace

FIGURE 21-15 Intensive diabetes treatment and cardiovascular disease in patients with type 1 diabetes—observations from the Diabetes Control and Complications Trial/Epidemiology of Diabetes Interventions and Complications (DCCT/EDIC) study. Cumulative incidence of the first of any of the predefined cardiovascular disease outcomes (A) and of the first occurrence of nonfatal myocardial infarction, stroke, or death from cardiovascular disease (B) . Compared with conventional treatment, intensive treatment reduced the risk of any predefined cardiovascular disease outcome by 42% (95% CI, 9%-63%; P=0.02) (A) and reduced the risk of the first occurrence of nonfatal myocardial infarction, stroke, or death from cardiovascular disease by 57% (95% CI, 12%-79%; P=0.02) (B) . (From Nathan DM, Cleary PA, Backlund JY, et al: Intensive diabetes treatment and cardiovascular disease in patients with type 1 diabetes. N Engl J Med 353:2643, 2005.)





discussed before, when lifestyle intervention and pharmacologic therapy fail to provide adequate control of diabetes in patients with severe obesity (BMI > 35 kg/m ²) and type 2 diabetes, bariatric surgery may be recommended and has been demonstrated to commonly resolve diabetes. ^{12,82} However, the long-term effect of bariatric surgery on diabetic CVD complications has not been adequately studied in randomized controlled trials.

Healthy Food Choices

In addition to the benefits of controlling calorie intake, certain food choices may help prevent CVD in patients with type 2 diabetes. 12,74 To prevent or to delay atherosclerosis, the ADA recommends that people with diabetes minimize their intake of saturated fat (< 7% of total calories), cholesterol (< 200 mg/day), and trans -fats and encourages the consumption of two or more servings of fish per week (not commercially fried) to provide omega-3 fatty acids. Diets high in fruits, vegetables, whole grains, and nuts and low in red meats, processed sweets, and sugar-containing beverages may also contribute to CVD risk reduction. Reduced intake of sodium lowers blood pressure (< 2300 mg sodium per day) and may also be beneficial for patients with symptomatic CHF (< 2000 mg sodium per day). Finally, moderate consumption of ethanol (less than one drink per day for women and two for men) may reduce CVD risk in patients with diabetes.

Physical Activity and Exercise

In addition to playing an important role in weight reduction and weight control, regular physical activity in patients with diabetes may independently lower the risk of cardiovascular complications. In several large cohorts of diabetic patients, higher levels of physical activity or aerobic fitness were associated with lower incidence of cardiovascular events and lower cardiovascular mortality, even with adjustment for glycemic control, BMI, and other risk factors. 124

The ADA recommends that people with diabetes perform at least 150 min/week of moderate-intensity aerobic physical activity (50% to 70% of maximum heart rate) as well as resistance training at least three times/week. ¹² Exercise programs should be tailored to individual patients, depending on age, prior levels of physical activity, risk of ischemic heart disease, and coexisting conditions.

Smoking Cessation

There is overwhelming evidence from both cross-sectional and prospective studies that smokers with diabetes have a higher risk of complications (both microvascular and macrovascular) and death compared with nonsmokers. ¹²⁵Diabetes and smoking act synergistically to increase cardio vascular risk, and the cardiovascular benefits of smoking cessation, especially in this high-risk population, cannot be overstated.

TABLE 21—5 Changes in Measures of Diabetes Control, Blood Pressure Control, Measures of Lipid/Lipoproteins Control, Albumin-to-Creatinine Ratio, and Prevalence of Metabolic Syndrome Among Participants in the Look AHEAD Trial at Year 1: Mean or Percent (Standard Error)

	Intensi Lifestyle Intervention	Diabetes Support and Education		
ve Measure	N = 2496	N = 2463	P -Value	
Baseline	86.5 (0.7)	86.5 (0.7)	0.93*	
Year 1	78.6 (0.8)	88.7 (0.6)	< 0.001*	
Change	- 7.8 (0.6)	2.2 (0.5)	< 0.001 ·	
Fasting Glucose (mg/dL)	454.0 (0.0)	452.0 (0.0)	0.24	
Baseline Year 1	151.9 (0.9) 130.4 (0.8)	153.6 (0.9) 146.4 (0.9)	0.21 · < 0.001 ·	
Change	- 21.5 (0.9)	- 7.2 (0.9)	< 0.001 ·	
Hemoglobin A1c (%)				
Baseline	7.25 (0.02)	7.29 (0.02)	0.26	
Year 1	6.61 (0.02)	7.15 (0.02)	< 0.001 ·	
Difference	- 0.64 (0.02)	- 0.14 (0.02)	< 0.001	
Danalina	75.2 (0.0)	70.7 (0.0)	0.22*	
Baseline	75.3 (0.9)	73.7 (0.9)	0.23*	
Year 1 Change	75.2 (0.9) - 0.1 (0.6)	75.9 (0.9) 2.2 (0.6)	0.54* 0.02 ·	
Baseline	128.2 (0.4)	129.4 (0.3)	0.01 ·	
Year 1	121.4 (0.4)	126.6 (0.4)	< 0.001 ·	
Change	- 6.8 (0.4)	- 2.8 (0.3)	< 0.001 ·	
Baseline Year 1	69.9 (0.2) 67.0 (0.2)	70.4 (0.2) 68.6 (0.2)	0.11 ·	
Change	- 3.0 (0.2)	- 1.8 (0.2)	< 0.001 · < 0.001 ·	
	0.0 (0.2)	(0.2)	1 0.00 1	
Baseline	49.4 (1.0)	48.4 (1.0)	0.52*	
Year 1	53.0 (1.0)	57.8 (1.0)	< 0.001*	
Change	3.7 (0.8)	9.4 (0.8)	< 0.001	
LDL-C (mg/dL)				
Baseline Year 1	112.2 (0.4) 107.0 (0.6)	112.4 (0.6)	0.78 · 0.74 ·	
Change	- 5.2 (0.6)	106.7 (0.7) - 5.7 (0.6)	0.49	
Change	- 3.2 (0.0)	- 3.7 (0.0)	0.10	
HDL-C (mg/dL) Baseline	43.5 (0.2)	43.6 (0.2)	0.80 ·	
Year 1	46.9 (0.3)	44.9 (0.2)	< 0.001	
Change	3.4 (0.2)	1.4 (0.1)	< 0.001	
Triglycerides (mg/dL)				
Baseline	182.8 (2.3)	180.0 (2.4)	0.38	
Year 1	152.5 (1.8)	165.4 (1.9)	< 0.001	
Change	- 30.3 (2.0)	- 14.6 (1.8)	< 0.001 ·	
Baseline	16.4 (0.7)	16.9 (0.8)	0.69 ·	
Year 1	18.4 (0.7)	15.4 (0.7)	0.005	
Change	- 3.9 (0.6)	- 1.5 (0.6)	0.002	
Metabolic Syndrome (%)				
Baseline	93.6 (0.5)	94.4 (0.5)	0.23	
Year 1	78.9 (0.8)	87.3 (0.7)	< 0.001	
Change	- 14.7 (0.8)	- 7.1 (0.7)	< 0.001	

^{*}Logistic regression with adjustment for clinical site. · Mantel-Haenszel test with adjustment for clinical site.

[^] Analysis of covariance with adjustment for clinical site.

From Pi-Sunyer X, Blackburn G, Brancati FL, et al: Reduction in weight and cardiovascular disease risk factors in individuals with type 2 diabetes: one-year results of the Look AHEAD trial. *Diabetes Care* 30:1374, 2007. Reproduced with permission.

Hypertension

Hypertension affects approximately 70% of patients with diabetes 46 and is more common in the elderly, in men, and in African Americans as well as in those with longer duration of diabetes, poorer glucose control, and concomitant protein uria. Hypertension adversely affects both microvascular and macrovascular disease risk diabetes group was twice as large compared to those without and has been estimated to account for between 35% and 40% of the diabetes at study entry. In the ALLHAT trial, the CVD effect of incremental CVD risk associated with diabetes.

Treatment of Hypertension in Patients with Diabetes Mellitus

change, and this most certainly applies to the prevention and treatment of hypertension. Frequent modest-intensity aerobic treatment proved superior for CVD outcomes. 130 Despite their exercise, alcohol and sodium moderation, and weight reduction all markedly contribute to improved blood pressure . Numerous should be considered among the evidence-based second-line classes of antihypertensive therapies have been proven safe and antihypertensive agents in diabetic patients, given their welleffective for the treatment of hypertension in patients with diabetes documented effects on blood pressure and CVD events. They on the basis of assessment of clinical outcomes in randomized trials are particularly suited in combination regimens with ACE by subanalyses of diabetes cohorts and also in trials exclusively inhibitors or ARBs, most of which are now available in fixedstudying patients with diabetes. A number of randomized trials dose combination formulations. have proven the efficacy of several classes of antihypertensive medications in this context, including angiotensin-converting blockers should be contraindicated for the treatment of patients enzyme (ACE) inhibitors, angiotensin receptor blockers (ARBs), calcium channel blockers, thiazide diuretics, and beta blockers. 126

Angiotensin-Converting Enzyme Inhibitors. Patients with diabetes and hypertension derive particular benefit from ACE inhibitors. ¹²⁶ Observations of treatment effects among patients with diabetes participating in long-term primary and second year blocker atenolol compared with placebo with regard to diabetescardiovascular risk prevention randomized trials with ACE related clinical outcomes among diabetic patients without prior inhibitors underscore the important clinical benefit of this class of drugs, 127-132 supporting their recommendation as the first-line captopril. 131 antihypertensive treatment of most patients with diabetes. 133 ACE inhibitors are also renoprotective, in both type 1 and type 2 diabetes. increased incidence of diabetes compared with nondiuretic

Angiotensin Receptor Blockers. Whereas the evidence base for ARBs is not as robust as that for the ACE inhibitors with regard to effects on CVD outcomes, the aggregate data support consideration metabolic differences between the beta blockers may be of of ARBs as an alternative treatment option in those patients intolerant of ACE inhibitors because of cough, rash, or angioedema. or metoprolol demonstrated in randomized head-to-head 126,133 ARBs may also be preferred for renoprotection in type 2 diabetes by a stronger evidence base from large randomized trials of patients with type 2 diabetes and impaired kidney function, such as IDNT and RENAAL. 135, 136 The ONTARGET randomized trial comparing ramipril versus telmisartan versus the combination of the two demonstrated comparable efficacy between telmisartan and ramipril, but it did not demonstrate incremental CVD benefits with the two drugs combined and in that setting increased adverse effects and intolerance. 137 Therefore, the combination of ACE inhibitors and ARBs is not routinely recommended.

Dihydropyridine Calcium Channel Blockers. Dihydro pyridine calcium channel blockers are well tolerated and effective at lowering blood pressure. Analyzes of diabetes subsets of randomized clinical trials have suggested CVD clinical benefits of similar or greater magnitude compared with those observed in the nondiabetic cohorts, including the evaluations of nitrendipine, nisoldipine, and Therapeutic Targets amlodipine. 126 In active controlled comparisons, amlodipine has Given the incremental CVD risk associated with hypertension in been proven superior to hydrochlorothiazide when it is added to a diabetes and the clearly demonstrated graded association background of benazepril therapy, 138 but in direct randomized between magnitude of blood pressure reduction and CVD comparisons of calcium channel blockers (fosinopril versus clinical risk reduction, patients with diabetes have been amlodipine, 139 enalapril versus nisoldipine 140), ACE inhibitors had identified as a special population warranting more aggressive superior efficacy on CVD outcomes.

affect glucose metabolism and have been associated with increased of risk for incident diabetes. However, despite

these glycometabolic effects, thiazides have been proven 363 effective at modifying cardiovascular risk among patients with diabetes and impaired fasting glucose concentration, I even in the setting of trials in which the thiazide was associated with increased diabetes incidence (eg, ALLHAT). 141 I The clinical efficacy of

thiazide diuretics among patients with diabetes is supported by the results of subanalyses of the 21 diabetic patients participating in the Systolic Hypertension in the Elderly Program (SHEP) trial, in which chlorthalidone was compared with placebo in patients aged > 60 years with isolated systolic hypertension. 142

In these analyses, the magnitude of treatment effect in the chlorthalidone was quantitatively similar to effects observed with amlodipine and lisinopril 127; and in the ADVANCE trial The keystone of cardiovascular risk prevention remains life style of high-risk diabetic patients that included the thiazide diuretic indapamide plus perindopril versus placebo, the combination possible effects on glucose metabolism, thiazide diuretics

> Beta blockers. For decades, common opinion was that beta with diabetes on the basis of concerns about adverse effects on glucose, insulin, and lipid metabolism, similar to observations with thiazide diuretics discussed before, as well as the masking of symptoms of hypoglycemia. Although these concerns persist, data from the UKPDS demonstrated superiority of the beta CVD, with treatment effects comparable to those observed with

> Subsequent trials, however, have demonstrated both antihypertensive therapies and slightly less efficacy at reducing major adverse CVD events compared with treatment with ACE inhibitors or calcium channel blockers. 143 In this context, importance, such as the superiority of carvedilol over ateno lol comparisons of glucose and lipid effects; the clinical relevance of these observations remains to be defined. 144,145

> Despite the accumulated clinical outcomes evidence, there continues to be some resistance across the broad clinical community to the use of beta blockers in patients with diabetes. However, whereas ACE inhibitors, calcium channel blockers, and thiazide diuretics appear to have some advantages over beta blockers, both metabolically and with regard to clinical outcomes, because achievement of aggressive blood pressure targets for patients with diabetes will commonly require three or four medications, beta blockers remain a therapeutic option in this setting. These agents should be considered part of the antihypertensive regimen in any diabetic patient with coexisting coronary artery disease, especially if MI has already occurred, and in the setting of systolic heart failure.

than usual blood pressure control. Targets for patients with Thiazide Diuretics. Thiazide diuretic medications adversely diabetes of < 130/80 mm Hg have been endorsed by a number

> **364** professional guidelines. 12,133,146 These recommendations, based largely on epidemiological data, have been supported

I by observations of more recent randomized clinical trials. In I the Hypertension Optimal Treatment trial, participants with I elevated diastolic blood pressure were randomized to treat ment to three different targets: < 90 mm Hg, < 85 mm Hg, and 21 < 80 mm Hg.

Whereas the overall trial failed to demonstrate significant differences by intensity of blood pressure treatment, a post hoc analysis of the subset of patients with diabetes exhibited a significant intensity-dependent reduction in CVD, including a 50% relative risk reduction for major adverse CVD events in the group with the lowest compared with the highest diastolic achieved in the diabetic subset of HOPE (139/77) and in the ADVANCE trials (136/73) provides further direct support for the safety and efficacy of such intensified blood pressure targets in the high-risk population of patients with diabetes. However, in the ACCORD randomized clinical trial, in which more than 10,000 patients with diabetes at increased CVD risk of glucose control to contemporary targets. were randomized to systolic blood pressure goals of < 120 mm nonfatal CVD events. 147a

Angiotensin-Converting Enzyme Inhibition Independent of Hypertension

treated with ACE inhibitors appears to extend beyond on the basis of the differences in blood pressure achieved. 126 These observations suggest pleiotropic benefits of ACE management of diabetic dyslipidemia. inhibition, including but not limited to favorable effects on glucose metabolism and insulin resistance. 148 This discordance between expected and observed effects of ACE inhibitors was first demonstrated in the Heart Outcomes Prevention Evaluation (HOPE) trial, 132 which randomized entry.

The results from HOPE demonstrated that treatment with ramipril titrated to 10 mg daily was associated with a 25% diovascular events and also reduced renal complications among the large subset of patients with diabetes enrolled in pressure between the groups of only 3 mm Hg. These the diabetes subset of the European trial on reduction of versus placebo in a cohort with a high prevalence of diabetes and in the ADVANCE trial comparing perindoprilwith only modest differences in blood pressure achieved ACE inhibitor use in all patients with diabetes at increased CVD risk, such as those aged > 40 years and those with CVD occurring at an earlier age, as recommended by contemporary professional guidelines. 12

Summary of Hypertension Management

Given the high prevalence and adverse microvascular and CVD consequences of hypertension in the diabetes population , aggressive control of blood pressure should be among the main therapeutic objectives in the management of this highrisk population of patients. Aggressive blood pressure management should have a target of < 130/80 mm Hg and, in addition to intensive lifestyle counseling, can use an arsenal of at least five classes of evidence-based medications for CVD risk modification. ACE inhibitors should be considered firstline therapy (and ARBs for those intolerant of ACE inhibitors), independent of blood pressure, for all diabetic patients aged > 55 years with additional cardiac risk factors or younger if prevalent CVD is present, especially in the context of their incremental benefit on renal outcomes.

Dyslipidemia

Dyslipidemia is common among patients with type 2 diabetes, especially in the majority of patients with significant insulin blood pressure target. 147 Likewise, the average blood pressure resistance, most commonly manifested as a characteristic pattern consisting of elevated triglyceride levels, decreased HDL levels, and only modest elevations of LDL but with an increased proportion of small, dense LDL particles. ^{62,64} The cornerstone for treatment of this dyslipidemia remains lifestyle interventions as described before, and lipid disturbances are also favorably affected by intensification

Several pharmacological agents are especially effective at Hg versus < 140 mg Hg, more intensive blood pressure control modifying this spectrum of abnormalities, including niacin and did not significantly reduce the combined risk of fatal or fibric acid derivatives, but the net CVD clinical effects of these interventions remain uncertain. In contrast, the statin drugs have a robust CVD clinical outcomes data base in the setting of diabetes and are the primary drugs advocated for use in patients with The reduction of CVD risk among patients with diabetes diabetes. Fish oil preparations also have an evidence base for favorable effects on CVD risk, although clearly much weaker than hypertension, with both microvascular and macrovascular that for statins, and can also be considered, given their beneficial benefits consistently greater in magnitude than that predicted effects on triglycerides. Several ongoing trials should help clarify the role of these agents alone and in combination with statins in the

Pharmacological Treatment of LDL-C

Despite the fact that marked elevations of LDL are not characteristic of diabetes dyslipidemia, statin medications have been firmly established as the primary lipid drug therapy to affect CVD risk patients with normal ventricular function and either diabetes modification. This is due to some degree to the propensity for with at least one additional risk factor or prevalent CVD to patients with diabetes to have small, dense LDL particles, resulting ramipril or placebo, independent of blood pressure at study in a much higher concentration of particles for any given mass of circulating LDL. This is reflected by increased LDL particle number and increased apolipoprotein B concentrations.

Prospective studies have demonstrated a strong, independent relative reduction in the long-term risk of major adverse car relationship between LDL particle size or concentration and the risk of coronary artery disease. 149,150 This is thought to be attributable to increased atherogenic propensity for the small, dense LDL particles that trial, despite just achieving a mean difference in blood by virtue of their ability to more readily cross into the arterial wall; and once there, they are more prone to oxidation to initiate and to observations have subsequently been supported by results in propagate the atheromatous process, to induce endothelial dysfunction, and to increase thromboxane formation. 151 On the cardiac events with perindopril in patients with stable basis of these observations and the demonstrated efficacy of the coronary artery disease (EUROPA) comparing perindopril approach, LDL-C remains the main therapeutic target in patients with diabetes.

The most convincing evidence for the efficacy of statins for indapamide combination versus placebo in patients with type primary CVD risk prevention in the setting of diabetes derives 2 diabetes. ACE inhibitor-based regimens reduced CVD risk, primarily from two randomized trials, 152,153 supported by a metaanalysis of statin use in diabetes. 154 In the Heart Protection Study between the groups (approximately 5 to 6 mm Hg). In (HPS) comparing simvastatin 40 mg daily versus placebo among a aggregate, the data reported to date support consideration of population of patients at increased CVD risk but not meeting contemporary indications for statin therapy, 5963 patients with diabetes were enrolled, including 67% without prior CVD and 41% with an LDL level below 116 mg/dL at study entry. 153 In the overall diabetes subset, simvastatin was associated with a 22% (P < 0.0001) relative reduction in the first occurrence of a major coronary event, stroke, or revascularization and a 20% reduction in coronary

subjects without prior CVD, major adverse CVD events were triglyceride-rich lipoproteins. 160 Despite these 21 lipid effects, decreased by 33% (P = 0.0003) with simvastatin.

152,2838 patients with type 2 diabetes and at least one other basis of the triglyceride and HDL effects. cardiovascular risk factor but without clinical indication for statin therapy were randomized to atorvastatin 10 mg versus placebo. In Trial (VA-HIT), 2531 subjects with a history of coronary artery this cohort with an average baseline LDL level of 120 mg/dL, disease and low HDL levels (< 40 mg/dL) were randomized to atorvastatin treatment resulted in a 37% (P = 0.001) relative treatment with gemfibrozil 1200 mg daily versus placebo. ¹⁶¹ In reduction in major cardiovascular events, prompting early a post hoc subgroup analysis of 391 participants with diabetes, termination of the study. In summary, these data support the gemfibrozil was associated with a 32% (P = 0.004) risk reduction recommendation that patients with diabetes at sufficient risk for in cardiovascular events, with a 41% (P = 0.02) reduction in cardiovascular events should be considered for statin therapy death attributable to coronary artery disease. 162 Key limitations regardless of baseline choles terol levels, underpinning the most of extrapolating the VA-HIT observations to contemporary recent guideline recommendations for statin use in all patients with practice include the small number of diabetes subjects enrolled, diabetes aged > 40 years or younger with prevalent CVD, 12 and with the exclusion of statin use during the trial, the exclusive an aggressive LDL target of < 70 mg/dL to be considered. 155

A number of other approved therapies for dyslipidemia interactions of gemfibrozil with statins, especially simvastatin. favorably affect LDL concentrations, including ezetimibe, bile acid binders, fibric acids, and niacin. However, none of these alternative in high-risk CVD cohorts have failed to demonstrate the agents affects LDL as potently as the statin medications do, and as superiority of fibrates versus placebo. In the Bezafibrate mentioned before, none has yet been proven through rigorous Infarction Prevention (BIP) trial, 3090 patients with coronary randomized trial assessment to consistently benefit CVD risk and artery disease and low HDL concentration were enrolled, outcomes. On that basis, these should be considered alternatives or including 309 with diabetes, and randomized to treatment with possibly add on therapies for those patients with diabetes intolerant bezafibrate 400 mg daily versus placebo. 163 After 6.2 years of of statins or unable to achieve therapeutic targets with statin average follow-up, bezafibrate failed to significantly reduce the monotherapy.

Pharmacological Treatment of Triglycerides and HDL

Elevation of triglyceride levels is a key component of diabetic dyslipidemia, and epidemiological observations have consistently demonstrated a robust independent association between high triglyceride levels and adverse CVD outcomes. Similarly, another hallmark feature of dyslipidemia associated with insulin resistance and diabetes is low HDL-C concentration, and across epidemiological studies, low HDL concentration is among the strongest and most consistent predictors of adverse CVD risk. However, results from randomized trials assessing the effects on CVD risk of various therapeutic strategies targeted at treating high triglyceride and low HDL levels have been discordant and have largely failed to prove the therapeutic concept.

Omega-3 Fatty Acids. Long-chain omega-3 fatty acids (fish oil) lower triglyceride levels on average by 50 to 75 mg/dL, dependent to some degree on baseline abnormality and dose of fish oil, with up to 30% reduction when doses of 4 g/day are used. Fish oil has minimal effects on HDL and total cholesterol, modestly raises LDL, and despite early concerns to the contrary, has no discernible effects on glycemic parameters in patients with diabetes. Complementing these net effects on lipid parameters, data from fibrates has failed to demonstrate clear CVD clinical benefit in randomized clinical outcomes trials including representation of patients with diabetes have consistently demonstrated improved car diovascular clinical outcomes. 157,158

Randomized clinical trials assessing the effects of these agents on cardiovascular outcomes in patients with diabetes are lacking. The most robust evidence to date is derived from a subanalysis of the Japan EPA Intervention Study (JELIS) of 4565 patients with it is used in combination and may increase the risk of statinimpaired fasting glucose or diabetes, demonstrating a 22% risk associated toxicities. reduction (P = 0.048) for major adverse CVD events with eicosapentaenoic acid treatment at 1.8 g daily. 159 Presently, a largescale randomized trial is under way, the Outcome Reduction with Initial Glargine Interven tion (ORIGIN) trial, that is assessing the CVD effects of fish oil versus placebo in a population of ~10,000 patients with impaired fasting glucose, impaired glucose tolerance, or early type 2 diabetes, including but not limited to those receiving 366 data for this class of drugs remain lacking. In addition, background statin therapy.

prime candidates for the treatment of diabetic dyslipidemia by development of sustained-release preparations of niacin. Niacin virtue of their favorable effects on a number of the I characteristic is also known to exacerbate insulin resistance, which may result disturbances. Fibrates lower triglyceride levels and modestly in increasing glucose levels in patients with type 21 2 diabetes. increase HDL concentration by inhibiting VLDL I production and

mortality (P = 0.02). Furthermore, in the cohort of 2912 diabetic activating lipoprotein lipase, which increases catabolism of results from randomized clinical trials have been mixed and have In the Collaborative Atorvastatin Diabetes Study (CARDS), largely failed to prove the cardiovascular benefit anticipated on the

> In the Veterans Affairs High-Density Lipoprotein Intervention enrollment of men, and the present awareness of adverse drug

Subsequently, three large-scale randomized trials of fibrates risk of sudden death or MI compared with placebo (13.6% versus 15.0%; P = 0.26); exploratory post hoc analyzes suggested benefit in subjects with triglyceride levels > 200 mg/dL as well as in patients with the metabolic syndrome, which should be interpreted as hypothesis-generating observations requiring confirmation. 163,164

In the Fenofibrate Intervention and Event Lowering in Diabetes (FIELD) trial, ^{165,9795} patients with type 2 diabetes not taking a statin at study entry were randomized to micronized fenofibrate 200 mg daily versus placebo. Despite an average of 5 years of follow-up and acquisition of 544 primary outcome events, fenofibrate failed to significantly reduce fatal and nonfatal MI risk (5.9% versus 5.2%; P = 0.16). This less than anticipated benefit was in part attributed to a significantly greater drop-in rate for nonstudy lipid-lowering drugs, predominantly statins, in the placebo group (17%) compared with the fenofibrate group (8%; P < 0.0001). The ACCORD trial also failed to demonstrate a CVD benefit of fenofibrate versus placebo in patients with type 2 diabetes at increased CVD risk who had residual elevation of triglycerides after achieving target LDL concentrations with statin therapy. 164a

In summary, randomized clinical outcome assessment of notable the context of contemporary prevalent use of statin medications with many results trending in the right direction but interpretation and generalizability substantially confounded by issues of concomitant statin use, both in the trials and in clinical practice. In addition, one fibrate, gemfibrozil, is known to increase circulating statin levels (especially simva statin) when

Niacin. Niacin is particularly attractive for the treatment of diabetic dyslipidemia as it increases HDL levels up to 35% and also reduces triglyceride levels 15% to 20%, reduces LDL concentration 10% to 15%, and increases LDL particle size. 166 Despite these consistently favorable lipid effects on the lipo protein spectrum of diabetic dyslipidemia, clinical outcomes clinical use is often confounded by the common side effect of I Fibric Acid Derivatives. Fibric acid derivatives (fibrates) 365 are flushing, which has been mitigated but not eliminated by the I

The only large-scale randomized trial reported to date

assessing the CVD effect of monotherapy with niacin is the Coronary Drug Project (CDP), 18 which randomized 1119 men with prior CVD to treatment with immediate-release niacin and compared CVD outcomes with placebo treatment in 2789 patients. Niacin therapy was associated with a modest 17% (P one of which included niacin.

relatively low numbers enrolled.

In addition to the uncertainties about the CVD efficacy of niacin monotherapy, few data are available regarding the incremental efficacy and safety of niacin added to statins. In the HDL-Atherosclerosis Treatment Study (HATS), 168,160 Insulin resistance and type 2 diabetes are associated with myriad also treated with antioxidants, suggesting an adverse of patients with diabetes. treatment interaction. Given the very small sample size,

National Heart, Lung, and Blood Institute, syndrome.

Therapeutic Targets

The key therapeutic target underpinning any treatment of diabetic dyslipidemia remains lifestyle intervention focused on optimization of medical nutrition therapy, weight control, and the habituation of adequate physical activity. The firstline pharmacologic therapy for diabetic dyslipidemia is the statin medications, as reflected in the most recent update of the NCEP Adult Treatment Panel (ATP III) 155 and supported by other professional society guidelines. 12,133,169 Whereas some minor differences in recommended lipid management strategies exist across the guidelines, those formulated by the ADA updated annually are the most current, 12 aged > 40 years or younger in the setting of either prevalent CVD or the presence of CVD risk factors other than diabetes.

The recommended LDL therapeutic targets are as follows: LDL < 100 mg/dL in the absence of underlying CVD; LDL < 70 mg/dL in the setting of prevalent CVD; and, at maximum tolerated statin dose, at least 30% to 40% reduction from baseline LDL concentration. Importantly, none of these rec ommendations for statin prescription is based on the baseline LDL level; instead, they are driven by the underlying CVD risk burden. The updated NCEP ATP III guidelines regard diabetes as a coronary disease equivalent and advocate an "optional" LDL goal of < 70 mg/dL in all patients with diabetes that is independent of underlying CVD prevalence or risk estimate. 155

Beyond LDL interventions, as reviewed in this section, little < 0.05) reduction in coronary death or nonfatal MI after 6 evidence is available to guide decision making with regard to other years and, after an extended 15-year post-trial follow-up lipid targets. For the treatment of patients who have persistently period, a 16% (P < 0.005) reduction in all-cause mortality. ¹⁶⁷ elevated triglyceride levels (> 200 mg/dL) after achieving However, interpretation of the CDP trial results is - therapeutic LDL targets, a consensus opinion across guidelines challenging, given an extreme rate of study drug advocates that the principal secondary lipid target should be nonnoncompliance in the niacin arm and comparisons of CVD HDL (ie, non-HDL = total choles terol - HDL), with target levels 30 events partitioned over six randomized treatment arms, only mg/dL higher than the individual patient's corresponding LDL target. This can be achieved by intensifying lifestyle intervention or Similar limitations of compliance and power apply to the statin prescription or by the addition of a second drug, such as fish accumulated data set regarding the CVD effects of niacin, oil, niacin, or a fibrate (but not gemfibrozil). As summarized in the largely limited by small studies and challenges of trial ADA guidance, however, the net safety and efficacy of add-on drug execution with excessive dropout among subjects taking therapy remain poorly understood, with a noted increase in the risk niacin. In addition, there has been limited power to assess for liver and muscle side effects. Despite the intuitive appeal of effects unique to patients with diabetes because of the treating triglyceride or HDL targets, such strategies lack a solid evidence base.

Antiplatelet Therapies

patients with low HDL concentration and obstructive abnormalities in platelet structure, life span, activation, and coronary disease were randomized to a combination of aggregation, yielding a prothrombotic state. ^{60,61} This is simvas tatin and niacin, simvastatin-niacin and antioxidant compounded by secondary platelet effects of other coagulation therapy, antioxidant therapy alone, or placebo. Compared factor abnormalities associated with diabetes, such as increased with placebo, simvastatin-niacin was associated with circulating levels of fibrinogen, thromboxane, plasminogen regression of coronary plaque and, on the basis of only 12 activator inhibitor 1, and von Willebrand factor, which contribute to versus 1 major adverse CVD events, statistically superior platelet activation and aggregation. ¹⁷⁰On that basis, the CVD effects clinical outcomes; these effects were eliminated in the group of antiplatelet therapies have been of great interest for the treatment

Aspirin. Whereas the evidence base for the use of aspirin to however, the validity of these observations is remarkably reduce CVD risk in the setting of prevalent CVD is well established, ¹⁷¹ the role of aspirin for primary CVD risk prevention is much less To more rigorously assess the CVD effects of niacin added well defined. 172 This is especially true in the setting of diabetes, for to statin therapy, a large ongoing study sponsored by the which the data are substantially limited by relatively few diabetic the patients studied to date. Analyzes from the diabetes subsets of Atherothrombosis Intervention in Metabolic Syndrome with reported clinical trials suggest an attenuation of aspirin benefit Low HDL/ High Triglycerides and Impact on Global Health among patients with diabetes, as demonstrated in the Outcomes (AIM-HIĞH), is comparing the effects of niacin Antithrombotic Trialists' meta-analysis of antiplatelet therapy added to simvastatin versus simvastatin alone on CVD (predominantly aspirin) comprising a patient mix of primary and outcomes in a randomized controlled trial of ~3300 patients secondary CVD risk and including assessment of almost 5000 with prevalent CVD, type 2 diabetes, and metabolic patients with diabetes. 171 That analysis revealed a significant relative odds reduction with antiplatelet therapy of 22% in the overall cohort but only a trend of 7% reduction in the subset with diabetes that was not statistically significant.

The Hypertension Optimal Treatment (HOT) randomized clinical trial evaluated the efficacy of aspirin 75 mg daily versus placebo on CVD event risk reduction among patients with hypertension, including 1501 (8%) with diabetes. 147 After a mean follow-up of 3.8 years, aspirin was associated with a 36% relative reduction in risk for MI and a 15% relative reduction in major cardiovascular events in the overall trial. The authors qualitatively commented that the results among the diabetes subset were similar, although no data were presented for these analyses.

Subsequently, the results from three randomized clinical trials of aspirin in the setting of primary CVD prevention have challenged recommending statin therapy for all patients with diabetes the utility of routine aspirin use for primary CVD prevention in diabetes. The Primary Prevention Project

(PPP) randomized trial assessed the CVD effects of 75 mg of aspirin and are the focus of numerous ongoing research programs. daily versus placebo and included 1031 (23%) patients with diabetes. 173 Treatment with aspirin was associated with a accumulated data, the routine use of aspirin at doses ranging from nonsignificant trend of 10% relative risk reduction for death, MI, 75 to 162 mg daily remains widely recommended by contemporary and cerebrovascular accident, contrasted with a significant 41% guidelines for primary CVD risk prevention for most adult patients reduction observed in the nondiabetic group. However, because of with diabetes, including those aged > 40 years or younger with early termination of the trial due to efficacy in the overall trial, only additional CVD risk factors. 12,133,183 The strength of such 42 primary events were evaluable within the diabetes subanalysis, recommendations is variable across guide lines, ranging from Level

thus markedly limiting the statistical power and validity of the stati modest, and the study was further limited by 50% withdrawal rate Diabetes mellitus is common and increasing globally, driven of study subjects by 5 years.

Aspirin for Diabetes (JPAD) trial randomized 2539 patients with prevent diabetes in the first place, and once it is present, continues type 2 diabetes to aspirin 81 to 100 mg daily versus placebo, with with efforts to mitigate the associated clinical risk, which is follow-up for a median of 4.4 years, accumulating 154 primary primarily CVD. Lifestyle intervention remains the cornerstone of events of cardiovascular death, MI, cerebro vascular accident, and such management; when it is applied effectively, it potently peripheral arterial disease. ¹⁷⁵ Treatment with aspirin was associated prevents diabetes and its myriad associated CVD risk factors. with a trend towards a 20% hazard reduction that did not achieve statistical significance (HR = 0.80; 95% CI, 0.58-1.10; P = 0.16).

risk prevention in diabetes, two large-scale randomized clinical strategies and glucose targets. Beyond glucose, interventions trials are underway. A Study of Cardiovascular Events in Diabetes broadly demonstrated to favorably affect CVD in the general (ASCEND) is designed to enroll 10,000 patients with type 1 or type population are especially effective for patients with diabetes, 2 diabetes without CVD in a double-blind, placebo-controlled trial including intensive blood pressure and LDL management added to with a factorial ran domization to treatment with (1) 100 mg aspirin lifestyle interventions. daily versus placebo and (2) omega-3 fatty acid 1 g daily versus placebo, with a primary endpoint of major adverse cardiovascular events (http://www.ctsu.ox.ac.uk/ascend/). The Aspirin and Simvastatin Combination for Cardiovascular Events Prevention -Trial in Diabetes (ACCEPT-D) plans to enroll 4700 patients with type 1 or type 2 diabetes to receive 100 mg aspirin plus simvastatin versus simvastatin alone in a prospective, open-label, blinded endpoint evaluation (PROBE) design trial to assess the cardiovascular efficacy of aspirin in primary prevention for patients with diabetes treated with statins. 176

In addition to overall uncertainty about the utility of aspirin in primary prevention populations with diabetes, additional uncertainty remains about the most appropriate dose of aspirin to prevent CVD in diabetes; most contemporary guidelines recommend doses of 75 to 162 mg/day. 12,133,177,178 However, all such aspirin dosing recommendations in the setting of type 2 diabetes are acknowledged to derive primarily from expert opinion, as randomized comparative data guiding the choice of daily aspirin dose for patients with type 2 diabetes are effectively nonexistent. It remains possible that higher doses of aspirin may be more effective in type 2 diabetes, as suggested by results from the Early Treatment Diabetic Retinopathy Study (ETDRS) ran dominated trial that demonstrated superior CVD outcomes with 650 mg daily of aspirin 10 versus placebo among patients with type 1 and type 2 diabetes, ¹⁷⁹ a concept that requires confirmation before clinical application.

Other Antiplatelet Therapies. Based on the platelet abnormalities of type 2 diabetes and the ongoing uncertainty with regard to aspirin described before, the possibility remains that the more potent antiplatelet agents or agents targeting novel therapeutic targets may be especially beneficial in 367 diabetes. Support for this concept derives from several published reports from acute coronary syndrome studies, demonstrating incremental efficacy of parenteral glycoprotein IIb/ I IIIa antagonists ¹⁸⁰ and of both currently available thienopyridine antiplatelet agents, clopidogrel and prasugrel. 181,182 These concepts remain unproven in the setting of primary 21 CVD risk prevention

In summary, despite the uncertainties deriving from the

primarily by the ever-increasing prevalence of type 2 diabetes . The The Japanese Primary Prevention of Atherosclerosis with management of this at-risk population begins with attempts to

Glucose control strategies, although effective for microvascular disease risk intervention, remain unproven with regard to CVD risk In the context of this uncertainty in the setting of primary CVD reduction, with ongoing uncertainty about the most appropriate

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CHAPTER 22

Metabolic Syndrome and Cardiovascular Disease

Shaista Malik and Nathan D. Wong

KEY POINTS

- The metabolic syndrome is a constellation of risk factors that even in the absence of diabetes precedes development of cardiovascular disease.
- Visceral obesity and ensuing insulin resistance have been proposed as central features of the pathophysiology of metabolic syndrome.
- The definitions of metabolic syndrome have evolved over time and include visceral obesity, dyslipidemia, hypertension, and hyperglycemia.
- The prevalence of metabolic syndrome is approximately 34% among U.S. adults and is growing.
- Initial evaluation of coronary heart disease risk in metabolic syndrome subjects without diabetes involves global risk estimation by Framingham or other algorithms for risk prediction.
- Novel risk factors such as highsensitivity C-reactive protein as well as subclinical atherosclerosis (from carotid ultrasound, computed tomography, or ankle-brachial index) can refine the estimation of cardiovascular disease risk.
- The treatment of metabolic syndrome centers around lifestyle modification, supplemented with appropriate pharmacologic management, when indicated, for dyslipidemia, hypertension, and hyperglycemia.

The metabolic syndrome is a clustering of risk including factors visceral hypertension, dyslipidemia, and hyperglycemia, each an important risk factor for the development of diabetes cardiovascular disease constellation of these risk factors was initially described by Reaven as syndrome X and included insulin resistance, hyperglycemia, hypertension, low high-density lipoprotein cholesterol (HDL-C), and high very-low-density lipo protein triglycerides. ¹ Crosssectional surveys indicate that in the United States, one third of adults and an alarming proportion of youth have the metabolic syndrome. 2,3

Although there have been advances in understanding the pathophysiology, epidemiology, and prognostic implications of the metabolic syndrome and treatment strategies for it, uncertainties persist about whether metabolic syndrome has utility beyond its individual components. Focus on the metabolic syndrome ensures that attention is drawn clearly to the risk of CVD at all levels of the health care system. In particular, the metabolic syndrome criteria with the basic screening tool of waist measurement allow a relatively simple stepwise approach with particular attention to early detection of those at risk so that intervention can start. Although the metabolic syndrome may influence the choice of drug therapies, its presence essentially denotes the need to emphasize lifestyle management in clinical practice. In this chapter, we outline the historical perspective, pathophysiology, and evolving definitions of the metabolic syndrome; its significance as a tool for cardiovascular risk assessment; and therapeutic options.

HISTORICAL PERSPECTIVES

As early as 1923, Kylin, a Swedish physician, described the clustering of cardiovascular risk factors, such as hypertension, obesity, and gout. ⁴ More recently, in 1988, Reaven linked insulin resistance to hyper glycemia and hypertension and called this clustering

"syndrome X." In time, syndrome X took on the name metabolic syndrome and similar synonyms, such as the insulin resistance syndrome and the cardiometabolic syndrome. The American Association of Clinical Endocrinologists in 2001 promoted the recognition of the metabolic syndrome as a diagnostic entity with its own ICD-9 code. This action gave physicians the ability to diagnose and to manage the syndrome as its own entity rather than as component diagnoses. Moreover, in the same year, the Third Adult Treatment Panel of the National Cholesterol Education Program promoted the utility of easily determined criteria for defining the metabolic syndrome, which remain the most widely used criteria in the United States. ⁵Similar criteria were proposed shortly thereafter by the International Diabetes Federation, except that they focused on abdominal obesity to be one of the three criteria required for diagnosis of the metabolic

Then, in a joint statement, the American Diabetes Association and the European Association for the Study of Diabetes (ADA-EASD) questioned the value of diagnosis of the syndrome. ⁷ The main concerns that the ADA-EASD joint statement outlined included ambiguous or incomplete criteria for the diagnosis of the metabolic syndrome, uncertain role of insulin resistance as the cause of the syndrome, CVD risk attributable to the metabolic syndrome possibly not being greater than that attributable to individual components of the syndrome, and treatment of the metabolic syndrome not being different from treatment of component features.

In contrast, the position of the American Heart Association and the National Heart, Lung, and Blood Institute is that recognition of the syndrome in clinical practice is encouraged for the identification of a multiplerisk-factor condition and to promote lifestyle therapies that will reduce all of the metabolic risk factors simultaneously." 8

These differing position statements are essentially matters of perspective. ^{9,10} In reviewing previous literature on the

372 metabolic syndrome, Blaha and Elasy found that some papers used the metabolic syndrome as a study exposure (the clinical and epidemiological perspective) and that others used it as an outcome (the pathophysiological perspective). In their analysis, they found that the ADA-EASD position statement aligns with the pathophysiological perspective and that the basis for 22 most of the expressed concerns is the imprecise definition of the syndrome and incomplete understanding of its pathophysiology. By contrast, the American Heart Association position statement aligns itself with the clinical epidemiologic perspective and finds the recognition of features of the metabolic syndrome of substantial clinical use in the identification of patients at high risk for atherosclerotic events.

PATHOPHYSIOLOGY

Insulin resistance was proposed by Reaven to play the causal role in the pathophysiology of metabolic syndrome. It is certainly closely related to several of the components of the metabolic syndrome, and several metabolic pathways linking insulin resistance to the other factors, such as dyslipidemia and hypertension, have been proposed. ^{11,12} Insulin is a hormone that facilitates glucose uptake in adipocytes, hepatocytes, and skeletal muscle (Fig. 22-1). It also regulates hepatic glucose production and lipolysis. Insulin resistance has been defined as a condition of decreased responsiveness of target tissues to normal levels of circulating insulin, resulting in hyperinsulinemia.

Insulin resistance arises from both genetic and acquired defects. A major contributor to the development of insulin resistance is an overabundance of circulating free fatty acids, released from an expanded adipose tissue mass. Free fatty acids reduce insulin sensitivity in muscle by inhibiting insulin mediated glucose uptake. Increased levels of circulating glucose increase pancreatic insulin secretion, resulting in hyperinsulinemia. In the liver, free fatty acids increase the production of glucose, triglycerides, and secretion of very-low-density lipoproteins. The consequence is the reduction in glucose transformation to glycogen and increased lipid accumulation in triglycerides.

Multiple mechanisms have been proposed to explain the link between hypertension and insulin resistance. Hyperinsulinemia is associated with adrenergic overactivity, leading to increased cardiac output and urinary catecholamine excretion. ¹³ Insulin also has an antinatriuretic effect, causing sodium retention and plasma volume expansion. Increased sympathetic activity also stimulates the renin-angiotensin system (RAS). There is also an effect of insulin resistance on endothelial function. The elevation in free fatty acids and tumor necrosis factor -a (TNF-a) and the decrease in adiponectin adversely affect endothelial function and promote atherogenesis. ^{14,15}

More recently, it has been suggested that low-grade inflammation underlies or exacerbates the syndrome. The excess adipose tissue observed in the metabolic syndrome results in overproduction of the inflammatory cytokines TNF -a , -interleukin-6, and C-reactive protein (CRP). $^{\rm 16}$

DEFINING METABOLIC SYNDROME

During the past decade, several different definitions of metabolic syndrome have been proposed and used (Table 22-1). This has led to confusion and lack of comparability between studies. The first attempt to define metabolic syndrome was by a World Health Organization (WHO) diabetes group in 1998, which proposed a working definition that could be modified as more information became available. ^{17,18} Most of the criteria in the WHO definition were based on Reaven's suggestions for syndrome X with the addition of obesity and microalbuminuria. The essential components of metabolic syndrome were considered to be glucose intolerance, impaired glucose tolerance or diabetes, and insulin resistance together with two or more of

the following: raised arterial pressure, raised triglycerides, or low levels of HDL-C; central obesity or body mass index (BMI) > 30 kg/m ²; and microalbuminuria (see Table 22-1). Other possible components of the syndrome, such as hyperuricemia, coagulation disorders, and raised circulating levels of plasminogen activator inhibitor 1 (PAI-1), were mentioned but not added as necessary components of the syndrome. The European Group for the Study of Insulin Resistance then produced a modification of the WHO criteria excluding people with diabetes and requiring hyperinsulinemia to be present. ¹⁹

In 2001, the US National Cholesterol Education Program (NCEP) Adult Treatment Panel III (ATP III) recognized the existence of the metabolic syndrome as a major contributor to cardiovascular risk. 5 A strong emphasis of the definition was on recognition of people at high risk for CVD in addition to the conventional risk factors of low-density lipoprotein cholesterol (LDL-C), smoking, and family history. They produced a set of criteria in which people had to meet three of the five criteria that were similar to those of the WHO group but also showed some significant differences. Specifically, these criteria included increased waist circumference, ele vated blood pressure, impaired fasting glucose, increased triglycerides, and low HDL-C. Central adiposity was represented by waist circumference. Also, HDL-C and raised triglycerides could count separately. Blood pressure was also slightly lower for the ATP III definition (130/85 mm Hg or higher) than for the WHO criteria, and NCEP ATP III restricted glucose to the fasting state and included known diabetes. It was generally agreed that the NCEP ATP III definition was simpler for use in clinical practice.

The International Diabetes Federation (IDF) believed there was a strong need for one practical definition that would be useful in any country for the identification of people at high risk for diabetes and CVD. 6 Central obesity, as assessed by waist circumference, was agreed as essential because of the strength of the evidence linking waist circumference with CVD and the other metabolic syndrome components; its inclusion as a required component meant that this could be used as an initial screening, followed by evaluation of the other components only if it is increased. The waist circumference cutoff selected was lower than the NCEP ATP III recommendations (above 94 cm in men and 80 cm in women), and ethnic-specific waist circumference cutoffs have been incorporated into the definition, including lower cutoffs in Asians/South Asians of 90 cm in men and 80 cm in women. These cutpoints are also recommended for those of Central and South American ancestry. The levels of the other variables were as described by ATP III, except that the revised cut point from the American Diabetes Association for impaired fasting glucose (100 mg/dL) was used. The consensus group also recommended additional criteria that should be part of further research into metabolic syndrome, including tomo graphic assessment of visceral adiposity and liver fat, bio markers of adipose tissue (adiponectin, leptin), apolipoprotein B, LDL particle size, formal measurement of insulin resistance and an oral glucose tolerance test, endothelial dysfunction, urinary albumin, inflammatory markers (CRP, TNF- a, interleukin-6), and thrombotic markers (PAI-1, fibrinogen).

Most recently, the NCEP ATP III definition has undergone a revision in the American Heart Association/National Heart, Lung, and Blood Institute scientific statement on the diagnosis and management of the metabolic syndrome ²⁰ (see Table 22-1). Many of these revisions bring ATP III in line with the IDF recommendations. Given the recent American Diabetes Association revision of the lower cut point of impaired fasting

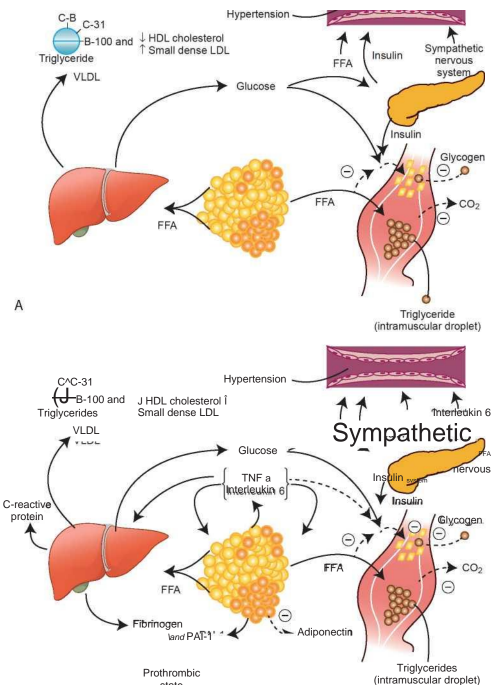


FIGURE 22-1 Pathophysiology of the metabolic syndrome (insulin resistance). A, Free fatty acids (FFA) are released in abundance from an expanded adipose tissue mass. In the liver, FFA increase production of glucose, triglycerides, and secretion of very-low-density lipoproteins (VLDL). Associated lipid and lipoprotein abnormalities include reductions in high-density lipoprotein (HDL) cholesterol and an increased density of low-density lipoproteins (LDL). FFA also reduce insulin sensitivity in muscle by inhibiting insulin-mediated glucose uptake. Associated defects include a reduction in glucose partitioning to glycogen and increased lipid accumulation in triglyceride (TG). Increases in circulating glucose and to some extent FFA increase pancreatic insulin secretion, resulting in hyperinsulinemia. Hyperinsulinemia may result in enhanced sodium reabsorption and increased sympathetic nervous system activity and contribute to the hypertension, as might increased levels of circulating FFA. B, Superimposed and contributory to the insulin resistance produced by excessive FFA is the paracrine and endocrine effect of the proinflammatory state. Produced by a variety of cells in adipose tissue including adipocytes and monocyte-derived macrophages, the enhanced secretion of interleukin-6 and tumor necrosis factor- a (TNF- a), among others, results in more insulin resistance and lipolysis of adipose tissue triglyceride stores to circulating FFA. Interleukin-6 and other cytokines are also increased in the circulation and may enhance hepatic glucose production, the production of VLDL by the liver, and insulin resistance in muscle. Cytokines and FFA also increases the production of fibrinogen and plasminogen activator inhibitor 1 (PAI-1) by the liver that complements the overproduction of PAI-1 by adipose tissue. This results in a prothrombotic state. Reductions in the production of the anti-inflammatory and insulin-sensitizing cytokine adiponectin are also associated with the metabolic syndrome. Lancet 365:1415, 2005.)

TABLE 22—1 Comparison of Definitions of Metabolic Syndrome

WHO (1999)

22

Diabetes or impaired fasting glycemia or impaired glucose tolerance or insulin resistance (euglycemic clamp: glucose uptake in the lowest 25%, hyperinsulinemia)

Plus two or more of the following: Obesity: BMI > 30 or waist-to-hip ratio of > 0.9 for men and > 0.85

Dyslipidemia: elevated triglycerides > 150 mg/dL (> 1.7 mmol/L) h HDL < 35 mg/dL (0.9 mmol/L) for men or < 39 mg/dL (1.0 mmol/L) for women Hypertension: blood pressure

> 140/90 mmHg Microalbuminuria: albumin excretion > 20 ^ g/min

EGIR (1999)

Insulin resistance, hyperinsulinemia: top 25% of fasting insulin values from non-diabetic population Plus two or more of the

following:
Central obesity: waist
circumference > 94 cm (hand)

or > 80 cm (woman)
Dyslipidemia: triglycerides

> 2.0 mmol/L (178 mg/dL) or HDL cholesterol

< 1.0 mmol/L (39 mg/dL) Hypertension: blood pressure

> 140/90 mm Hg or above antihypertensive

Fasting plasma glucose > 6.1 mmol/L (110 mg/dL) but no diabetes

NCEP ATP III (2001), AHA/NHLBI (2005)

Three or more of the following: Central obesity: waist circumference

> 102 cm (40 inches) in men, > 88 cm (35 inches) in women

Hypertriglyceridemia: triglycerides > 1.7 mmol/L (150 mg/dL) or *medication for elevated triglycerides

Low HDL-C: < 1.03 mmol/L (40 mg/dL) for men, < 1.29 mmol/L (50 mg/dL) for

< 1.29 mmol/L (50 mg/dL) fo women, *or medication for low HDL-C

Hypertension: blood pressure > 130/85 mm Hg or above *antihypertensive medication

*Fasting plasma glucose > 5.6 mmol/L (100 mg/dL) or on medication for hyperglycemia

IDF (2005)

Central obesity (ethnicity specific), defined by waist circumference: Europids: > 94 cm (37 inches) in men and > 80 cm (31.5 inches) in women

Asians/South Asians: > 90 cm in men or > 80 cm in women Plus two or more of the following: Hypertriglyceridemia: triglycerides > 1.7 mmol/L (150 mg/dL)

Low HDL-C: < 1.03 mmol/L (40 mg/dL) for men, < 1.29 mmol/L (50 mg/dL) for women

Hypertension: blood pressure > 130/85 mm Hg or above antihypertensive medication Fasting plasma glucose

> 5.6 mmol/L (100 mg/dL)

glucose to 100 mg/dL, this lower cut point has now been adopted in the revised definition. In addition, the criteria for elevated blood pressure, elevated triglycerides, and low HDL-C now also include medication for these conditions. The statement also comments on possible ethnic differences and that lower waist values may be adopted in certain ethnic groups, such as those recommended by the IDF.

PREVALENCE AND EPIDEMIOLOGY

The most recent data from the National Health and Nutrition Examination Survey (NHANES) 2003-2006 show a prevalence of metabolic syndrome of 34% among US adults by the NCEP ATP III guidelines criteria. ²¹ This estimate is identical to that with use of data from NHANES 1999-2002, which showed a prevalence (age adjusted) of metabolic syndrome of 34.4% among men and 34.5% among women by NCEP definition, with higher estimates of 40.7% and 37.1%, respectively, by the IDF definition ²² (Fig. 22-2). Earlier data among US adults show a substantially lower prevalence of approximately 25% in US adults examined in 1988-1994, ²³ with increases in prevalence from 1988-1994 to 1999-2000 noted to be most dramatic among women (increase of 23.5% during this period) and attributable mainly to increases in high blood pressure, waist circumference, and hypertriglyceridemia.

Wide variations in the prevalence of metabolic syndrome are observed across ethnic groups within the United States and worldwide but depend on the definition used. Despite attempts in recent years to reach an agreement on the definition of the metabolic syndrome, comparison of prevalences published for different populations is difficult because studies often differ with respect to the study design, the sample selection, the year that a study was conducted, the precise definition of the metabolic syndrome used, and the age and sex distribution of the population itself. Despite these obstacles, Cameron and associates 25 reported on the prevalence of the NCEP ATP III definition of the metabolic syndrome among various populations around the world (Fig. 22-3). Looking at those studies that include a population sample aged 20 to 25 years and older, the prevalence varies from 8% (India) to 24% (United States) in men and from 7% (France) to 46% (India) in women.

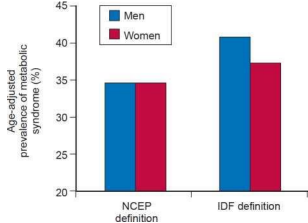


FIGURE 22-2 Prevalence of metabolic syndrome by National Cholesterol Education Program (NCEP) and International Diabetes Federation (IDF) definitions, US adults, National Health and Nutrition Examination Survey 1999-2002. (Modified from Ford ES: Prevalence of the metabolic syndrome defined by the International Diabetes Federation among adults in the US Diabetes Care 28:2745, 2005.)

In a large United Kingdom study, South Asians had the highest prevalence of metabolic syndrome (29% in men and 32% in women by the NCEP definition) and European women the lowest (14%). ²⁶ In a large study involving 11 European cohorts, prevalence with use of a modified WHO definition was slightly higher in men (15.7%) than in women (14.2%). ²⁷ Also, in Greek adults, age-adjusted prevalences of metabolic syndrome were 24.5% by the NCEP ATP III definition versus 43.4% by the IDF definition. ²⁸ Lower prevalence rates were recently noted by the ATP III definition among 2100 Italian adults: 18% in women and 15% in men. ²⁹

Ethnic-specific data among US adults have shown metabolic syndrome to be most prevalent among Mexican Americans; among African Americans, in particular men, prevalence was lower than in whites 22 (Fig. 22-4). From the Mexico City Diabetes Study, prevalence estimates from 1997-1999 show 39.9% of men and 59.9% of women to have the metabolic

^{*}Revisions incorporated by the American Heart Association/National Heart, Lung, and Blood Institute (AHA/NHLBI) definition (2005). EGIR, European Group for the Study of Insulin Resistance; IDF, International Diabetes Federation; NCEP ATP III, National Cholesterol Education Program Adult Treatment Panel III; WHO, World Health Organization.

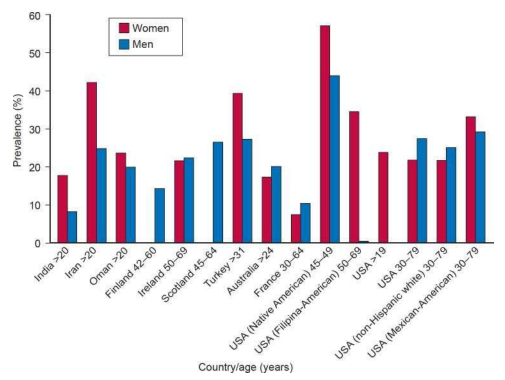


FIGURE 22-3 Worldwide prevalence of metabolic syndrome by NCEP ATP III definition. (Modified from Cameron AJ, Shaw JE, Zimmet PZ: The metabolic syndrome: prevalence in worldwide populations. Endocrinol Metab Clin North Am 33:351, 2004.)

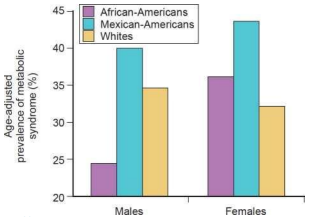


FIGURE 22-4 Prevalence of metabolic syndrome by ethnicity and gender, US adults, National Health and Nutrition Examination Survey 1999-2002. (Modified from Ford ES: Prevalence of the metabolic syndrome defined by the International Diabetes Federation among adults in the US Diabetes Care 28:2745, 2005.)

syndrome, representing little change from 1990-1992 among men (38.9%) but a decrease in women from that period (65.4%). Increases in prevalence were attributed to those with elevated waist circumference, elevated fasting glucose value, and low levels of HDL-C. ³⁰ These data, however, contrast with a nationwide study in Mexico involving 2158 men and women aged 20 to 69 years, in which the age-adjusted prevalence of metabolic syndrome was noted to be 13.6% by WHO criteria and 26.6% by the NCEP ATP III definition (and 9.2% and 21.4%, respectively, among those without diabetes). ³¹

Among Asian populations, lower prevalences of metabolic syndrome are generally noted. For example, among Hong Kong Chinese, prevalence was 9.6% by the NCEP definition and 13.4% by the WHO definition, ³² but in another study of Hong Kong

Chinese, prevalences were greater: 16.7% by the NCEP definition but 21.2% with incorporation of lower waist circumference criteria recommended for Asians by the WHO (> 80 cm for men and > 90 cm for women). 33 In 1230 Korean adults aged 30 to 79 years, the prevalence by WHO criteria was 21.8% in men and 19.4% in women; however, this increased to 34.2% of men and 38.7% of women with use of the modified NCEP definition. 34 Japanese and Mongolian adults had prevalences of only 6% and 12%, respectively, by ATP III criteria; however, these estimates did not factor in lower IDF-recommended waist circumference cut points for Asians. 35

Âmong US adolescents aged 12 to 19 years from the NHANES 2001-2006 survey, a prevalence of metabolic syndrome of 8.6% overall (10.8% in males and 6.1% in females) has been recently reported, ³⁶ a clear increase from an overall prevalence of 6.4% in 1999-2000 and 4.2% in 1988-1992. ³⁷ In a school-based cross-sectional study of 1513 black, white, and Hispanic teens, the overall prevalence of NCEP-defined metabolic syndrome was 4.2%, and that of WHO-defined metabolic syndrome was 8.4%; among obese teenagers, this increased to 19.5% and 38.9%, respectively. Moreover, non- white teens were more likely to have metabolic syndrome defined by WHO criteria. ³⁸ A recent review on the prevalence of metabolic syndrome in children and adolescents found ranges from 1.2% to 22.6%, with rates of up to 60% observed in the overweight and obese, in 36 studies from general population and community-based sampling. ³⁹

CARDIOVASCULAR RISK IN PERSONS WITH THE METABOLIC SYNDROME Prediction of Diabetes

Recent analysis of data from the Prospective Study of Pravas tatin in the Elderly at Risk (PROSPER) study and the British

376 Regional Heart Study (BRHS) found a strong association of metabolic syndrome with the incidence of type 2 diabetes. 40 I Metabolic syndrome predicted an increased risk of diabetes in women a twofold greater risk of mortality from CHD and CVD in PROSPER participants (HR = 4.41; 95% CI, 3.33-5.84), and I an even persons with metabolic syndrome; even those with metabolic stronger association was observed in BRHS participants (HR = 7.47; syndrome but without diabetes and those with only one or two 4.90-11.46). Similarly, data from the San 22 Antonio Heart Study (n metabolic syndrome risk factors were at an increased risk of death = 2559) showed that the metabolic syndrome predicted diabetes from CHD and CVD. Increased risks associated with metabolic beyond glucose intolerance alone. 41

Prediction of Cardiovascular Events and Mortality

The risk of CVD has been well studied and documented in persons with diabetes; in fact, diabetes is considered to be a coronary heart disease (CHD) risk equivalent according to the NCEP ATP III guidelines. 5 The East-West study showed that in the absence of prior myocardial infarction, persons with diabetes have a risk of future cardiovascular mortality similar to that of persons with a prior myocardial infarction without diabetes. The presence of both diabetes and prior myocardial infarction is associated with an even higher risk of cardiovascular mortality. 42 In our study involving up to 12 years of follow-up of 6255 adults from NHANES 1988-1994 we demonstrated total mortality to be similar among persons with diabetes but without preexisting CVD and those with CVD without diabetes 43 (Fig. 22-5).

The combination of diabetes and metabolic syndrome is associated with a much higher prevalence of CHD, and even those with metabolic syndrome in the absence of diabetes have a higher prevalence of CHD than do those with diabetes who metabolic syndrome is highly prevalent. Among a crosssectional survey of 1117 patients with CHD, cerebro vascular disease, peripheral vascular disease, or abdominal aortic aneurysm, an overall prevalence of metabolic syndrome was noted to be 46%; it was 58% in those with peripheral vascular disease, 41% in those with CHD, 43% in those with CVD, and 47% in those with abdominal aortic aneurysm. Moreover, age

did not have an impact on these prevalences. 45

We recently demonstrated in the US population of men and syndrome held similarly for men and women. Moreover, those with diabetes had a risk of future mortality similar to that of those with preexisting CVD. Those with both diabetes and preexisting CVD had the highest risk. 43 These observations are consistent with other reports documenting the prognostic importance of the metabolic syndrome; among 6447 men in the West of Scotland study, it predicted both incident diabetes (HR, 3.50; 95% CI, 2.51 4.90) and CHD events (HR = 1.30; 95% CI, 1.00-1.67) 46; and among 1209 Finnish men observed for 11.4 years, it predicted increased CVD mortality (2.6- to 4.2-fold increased risk, depending on definition used) and total mortality (1.9- to 3.3-fold increased risk, depending on definition used). 47

More recently, Jeppesen and colleagues 48 presented an analysis of data from a large Danish population-based study of 2493 men and women and used both insulin resistance and metabolic syndrome as predictors of incident CVD. They reported that both insulin resistance and metabolic syndrome were independent predictors; the relative risk for CVD was 1.49 (95% CI = 1.07-2.07) for insulin resistance as quantified by homeostasis model assessment (HOMA-IR) and 1.56 for NCEP-defined metabolic syndrome (95% CI = 1.12-

Other US population-based studies have also demonstrated a relationship of metabolic syndrome to cardiovascular event risk. In the Framingham Offspring Study, Rutter and coworkers 49 showed do not have the metabolic syndrome. ⁴⁴ Conversely, among an age-, sex-, and CRP-adjusted hazard ratio for metabolic syndrome those with preexisting atherosclerotic vascular disease, of 1.8 (95% CI = 1.4-2.5) for prediction of incident CVD events during 7 years. Moreover, among 12,089 black and white middle-aged individuals in

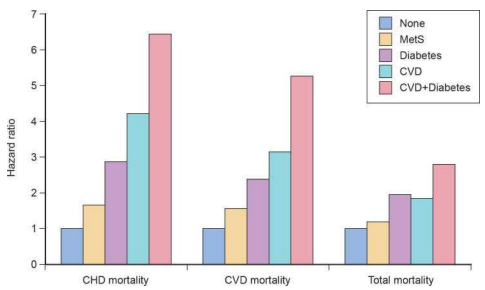


FIGURE 22-5 Cardiovascular disease and total mortality in U.S. men and women aged 30 to 74 years: age-, gender-, and risk factor-adjusted Cox regression, NHANES II follow-up (n = 6255). In comparison to those without metabolic syndrome, diabetes, or CVD, metabolic syndrome: P < 0.05 for CHD mortality and P < 0.01 for CVD mortality; diabetes, CVD, and CVD plus diabetes: P < 0.010.001 for CHD, CVD, and total mortality. CHD, coronary heart disease; CVD, cardiovascular disease; MetS, metabolic syndrome. (Modified from Malik S, Wong ND, Franklin SS, et al: Impact of the metabolic syndrome on mortality from coronary heart disease, cardiovascular disease, and all causes in United States adults. Circulation 110:1239. 2004.)

the Atherosclerosis Risk in Communities (ARIC) study, in which Stroke Prediction metabolic syndrome was found prevalent in 23% of those without diabetes or prevalent CVD at baseline, during an average 11 years of follow-up, those with versus without metabolic syndrome were 1.5 to 2 times more likely to develop CHD in risk factor-adjusted syndrome, those with metabolic syndrome but without diabetes had analyses. However, met abolic syndrome did not improve risk prediction beyond that achieved by the Framingham risk score. ⁵⁰

In 2175 elderly subjects in the Cardiovascular Health Study, metabolic syndrome defined by the ATP III but not by the WHO criteria was associated with a significant 38% increased risk (HR, 1.38; 95% CI = 1.06-1.79) of coronary or cerebrovascular events. 51 In the San Antonio Heart Study, among 2815 subjects aged 24 to 64 years, the NCEP metabolic syndrome definition predicted all-cause mortality (multivariable hazard ratio = 1.47; 95% CI = 1.13-1.92), but the WHO metabolic syndrome definition did not; among those without diabetes or prior CVD, the NCEP metabolic syndrome metabolic syndrome was associated with a 2.6-fold greater definition predicted only cardiovascular mortality (HR, 2.01; 95% CI = 1.13-3.57); there was also evidence for stronger relationships of metabolic syndrome with cardiovascular mortality in women Metabolic Syndrome Risks Among Subjects compared with men. 52 Among a large, primarily healthy cohort of 19,223 men who received a clinical examination and fitness examination, adjusted relative risks for all-cause and cardiovascular Among subjects with established CVD, the metabolic syndrome mortality were 1.29 (1.05-1.57) and 1.89 (1.36-2.60), respectively, is also associated with future CVD event risk. Among subjects among those with versus without metabolic syndrome. 53 Additional with acute coronary syndromes within the Myocar dial adjustment for cardiorespiratory fitness, however, resulted in Ischemia Reduction with Aggressive Cholesterol Lowering associations being no longer significant. Also, among 10,950 men in (MIRACL) trial, 38% of patients met the criteria for metabolic the Multiple Risk Factor Intervention Trial (MRFIT), modified syndrome; those with metabolic syndrome had a hazard ratio NCEP-defined metabolic syndrome was associated with increased of 1.49 (95% CI, 1.24-1.79) for the primary end point of death, hazard ratios during a median of 18.4 years of follow-up for total nonfatal myocardial infarction, cardiac arrest, or recurrent mortality (1.21 [1.13-1.29]), CVD mortality (1.49 [1.35-1.64]), and unstable myocardial ischemia. 62 Within the GISSI-Prevenzione CHD mortality (1.52 [1.34-1.70]), with elevated blood glucose and trial, among 11,232 patients with a prior myocardial infarction, low HDL-C being the most predictive factors of CVD mortality those with metabolic syndrome had a 29% greater risk of death among those men with metabolic syndrome. 54

diabetes, Ford 55 noted that among studies that used the exact NCEP 63 Finally, of interest are data from the Scandinavian definition of the metabolic syndrome, relative risks (and 95% Simvastatin Survival Study (4S) showing, among 3933 confidence intervals) associated with the metabolic syndrome were nondiabetic subjects with known CHD, those with the 1.27 (0.90-1.78) for all-cause mortality , 1.65 (1.38-1.99) for CVD, and metabolic syndrome to have at least as great (if not greater) 2.99 (1.96-4.57) for dia betes. For the WHO definition, corresponding reduction in the risk of total mortality (RR, 0.54), coronary estimates were 1.37 (1.09-1.74), 1.93 (1.39-2.67), and 2.60 (1.55-4.38). mortality (RR, 0.39), or major coronary artery disease events The authors concluded population attributable fractions of the (RR, 0.59) as those without the metabolic syndrome (0.72, 0.62, metabolic syndrome to be 6% to 7% for all-cause mortality, 12% to and 0.71, respectively). 64 17% for CVD, and 30% to 52% for diabetes.

The association between metabolic syndrome and CVD events Global Risk Assessment of Metabolic may be attenuated in certain population subgroups. A study by Syndrome Sattar and coworkers ⁵⁶ fueled the controversy over the importance

To best target treatment strategies, adequate assessment of risk of metabolic syndrome for determining vascular risk. A nonsignificant relationship was observed between metabolic syndrome and incident CVD in an elderly cohort from the PROSPER study (HR, 1.07; 95% CI = 0.86 1.32), and a weak association was observed in a similar cohort from the BRHS study (HR, 1.27; 1.04-1.56). In another study looking at the predictive value of metabolic syndrome in the elderly, Mozaffarian and colleagues 57 used data from the Cardiovascular Health Study and examined data from 4258 US adults. Although those with metabolic syndrome had a 22% higher mortality risk (RR, 1.22; 95% CI, 1.11-1.34), in looking at the population attributable risk fraction [PAR %], higher proportions of death were attributable to elevated fasting glucose and hypertension (PAR, 22.2%) than to metabolic syndrome (PAR, 6.3%). This study reinforces the concept that there is limited short-term risk assessment value in using metabolic syndrome. In general, the metabolic syndrome may help identify younger cohorts who face a high longterm cardiovascular risk.

Finally, a large meta-analysis of the risk of incident cardiovascular events and death associated with metabolic syndrome analyzed data from 37 studies and 172,573 individuals. Metabolic syndrome in this analysis had a much stronger association with cardiovascular events and death, with a relative risk of 1.78 (95% CI = 1.58-2.00) (Fig. 22-6). This relationship was stronger in women and remained significant after 22 adjustment for traditional cardiovascular risk factors. 58

odds of stroke. 61

a 1.49-fold greater odds for ischemic stroke or transient ischemic attacks (95% CI, 1.20-1.84), whereas those with diabetes had a 2.29fold increased odds (95% CI, 1.88-2.78); risks were higher in women than in men. ⁵⁹ Case-control studies have also recently reported on the association of metabolic syndrome with stroke. In a Japanese study, among 197 stroke survivors and 356 matched controls, metabolic syndrome was associated with a significant 3.1-fold greater odds of stroke. 60 In another case-control study in Greece involving 163 stroke survivors aged 70 years and older and 166 controls, in risk factor-adjusted analyses,

Metabolic syndrome is also related to the risk of stroke. Among

14,284 subjects with CHD, 26% of whom had the metabolic

with Known Cardiovascular Disease

and 23% greater risk of major cardiovascular events; these risks In a meta-analysis of risks for all-cause mortality, CVD, and were amplified in diabetic patients (68% and 47%, respectively).

for CVD is needed in persons with metabolic syndrome. Initial evaluation of risk can be determined by Framingham risk scores, 65 given the significant heterogeneity in estimated risk of persons with metabolic syndrome. In a study applying Framingham global risk algorithms to the US population with metabolic syndrome, 38.5% were classified as low risk (< 6% 10-year risk of CHD), 8.5% were classified as moderate risk (6% to 10% 10-year risk of CHD), 15.8% were classified as moderately high risk (10% to 20% 10-year risk of CHD), and 37.3% were classified as high risk (> 20% 10-year risk of CHD, or pre-existing CVD or diabetes). 66

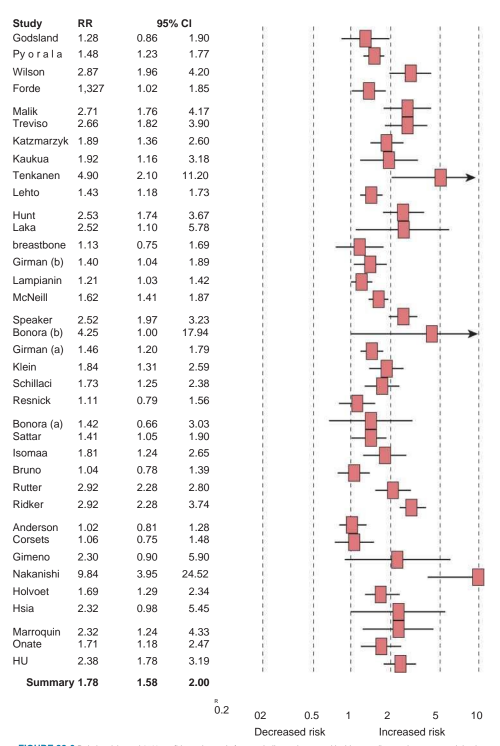


FIGURE 22-6 Relative risks and 95% confidence intervals for metabolic syndrome and incident cardiovascular events and death. Studies are listed in chronological order by the year in which their cohorts were created (except for the last study listed, which includes multiple cohorts). Results are for available analyzes of incident cardiovascular disease and death and may differ from the results of the total study populations. Boxes represent the relative risk, and lines represent the 95% confidence interval for studies. The diamond represents the pooled relative risk, and its width represents its 95% confidence interval. (Modified from Gami AS, Witt BJ, Howard DE, et al: Metabolic syndrome and risk of incident cardiovascular events and death. J Am Coll Cardiol 49:403, 2007.)

cholesterol or increased LDL-C levels, even if only minimal metabolic syndrome in predicting CVD events. 74 In addition, the elevations of defined metabolic syndrome risk factors are present, authors have reported that in the NHANES 1999-2000 sample, may be at intermediate or higher risk of CHD. However, an those with increased hsCRP levels and metabolic syndrome had a important limitation of Framingham risk or other global risk similar odds of CVD as those with diabetes and low hsCRP levels, algorithms is that they often do not include critical metabolic and those with diabetes and high hsCRP levels had the highest syndrome risk factors such as fasting glucose concentration or odds of CVD. 75 elevated triglycerides, which, although possibly not providing additive predictive value in a general population, could be critically hsCRP can identify within the primary prevention setting a important in stratifying risk in those with metabolic syndrome. subset of patients (those with elevated hsCRP levels) most likely Therefore, in situ ations in which a calculated global risk score to benefit from preventive therapy with a statin. Both patients results in a borderline figure (eg, 18% to 19% 10-year risk), the with and without metabolic syndrome, all of whom had hsCRP presence of significant metabolic risk factors not included in the levels > 2 mg/L, benefited from treatment. ⁷⁶ Disputing the global risk algorithm may warrant the individual to be stratified to relative importance of hsCRP, a recent comprehensive metaa higher risk stratum (eg, > 20% or CHD risk equivalent status in analysis from the Emerging Risk Factors Collaboration this case). The scientific statement on the clinical management of the examined data from 54 studies and 160,309 participants and metabolic syndrome released by the American Heart Association found that and National Heart, Lung, and Blood Institute noted, however, that cardiovascular risk factors resulted in attenuation of the linear the Framingham algo rithms do capture most of the risk for CVD in relationship between hsCRP concentration and CHD, stroke, persons with the metabolic syndrome and that adding obesity, and other vascular mortality. 77 The role of other novel risk triglyceride levels, and fasting glucose concentration does not markers, such as fibrinogen, interleukin (IL-1, IL-6), and appear to increase the power of prediction. 67

Diabetes Study (UKPDS) in 2001 developed a risk engine based on documented before any recommendations can be made. data from 4540 male and female patients with diabetes to predict the risk of new CHD events. Unlike previously published equations, Screening for Subclinical Atherosclerosis this model is diabetes specific and incorporates glycemia, systolic blood pressure, and lipid levels in addition to age, sex, ethnic group, smoking status, and time since diagnosis of diabetes. 68 Recent reports have examined the performance of this risk engine in relation to the Joint British Societies (JBS) risk calculator 69 and the earlier version of the Framingham risk equations that incorporated diabetes status. 70 Among 700 patients with type 2 diabetes, the of CHD 78,79 whereby those found to have clinically significant UKPDS risk calculator identified a higher mean 10-year CHD risk (21.5%) than the JBS risk calculator did (18.3%). 71

The more recent report compared the ability of the UKPDS risk engine and the Framingham risk equation to predict events that actually occurred among 428 subjects with newly diagnosed type 2 diabetes observed for a median of 4.2 years. The Framingham risk equations significantly underestimated the overall number of cardiovascular events by 33% and coronary events by 32%, compared with a lower and nonsignificant underestimation of normally recommended for those with diabetes. coronary artery disease events of 13% by the UKPDS risk engine, although both similarly performed in terms of discrimination and calibration for a 15% 10-year CHD risk threshold. 72 Another risk calculator was derived from the Prospective Cardiovascular Munster (PROCAM) study. In another study, adjustment of the PROCAM estimated global risk to include BMI or waist circumference corresponded very well with cardiovascular event rates. 73

Utility of Novel Biomarkers for Additional Risk Assessment in the Metabolic Syndrome

Once global risk assessment is done, additional information about the CHD risk for a given individual can be made with information obtained from novel biomarkers. For example, there has been Carotid Ultrasound interest in whether the addition of such risk factors as high- Intima-media thickness (IMT) of carotid arteries, as assessed sensitivity CRP (hsCRP), fibrinogen, and small dense LDL will noninvasively by carotid ultrasonography, is also a useful further add to the prediction of risk in persons with metabolic measure of preclinical atherosclerosis. Carotid IMT has been syndrome. It has been shown that hsCRP levels add predictive value for CVD risk among individuals with metabolic syndrome. In the Nurses' Health Study,

among persons with metabolic syndrome, age-adjusted incidence rates of future CVD events of 3.4 and 5.0 per 1000 person-years were demonstrated for those with hsCRP levels I of 3 mg/ L or more and levels of less than 3 mg/L, respectively, with additive effects of higher hsCRP levels also I observed for those with four or five metabolic syndrome risk factors. Framingham investigators

Older persons or those who are smokers or have increased total recently reported hsCRP 22 levels to provide additive value over

Importantly, the JUPITER trial has shown that screening for statistical adjustment for conventional adiponectin levels, in providing additive risk strati fication in For those with diabetes, the United Kingdom Prospective persons with metabolic syndrome needs to be examined and

Evaluation of subclinical atherosclerosis may have important implications for persons with metabolic syndrome, given the uncertainty of risk assessment on the basis of global risk assessment alone. Given recent recommendations to target atherosclerosis screening for those with intermediate global risk atherosclerosis could have their risk level stratified upward (eg, reclassification of an intermediate-risk individual as high risk), such screening may have implications for refining risk assessment in many persons with metabolic syndrome. Although such screening in persons with diabetes could also offer improved risk stratification, as diabetes is considered a CHD risk equivalent and aggressive treatment guidelines already exist for those with diabetes, such evaluation is not

Ingelsson and colleagues 80 evaluated the incidence of CVD associated with metabolic syndrome and diabetes according to the presence or absence of subclinical disease, which was categorized on the basis of any abnormalities on carotid ultra sound or ankle-brachial blood pressure, left ventricular hypertrophy on echocardiography or electrocardiography, or observed abnormal urinary albumin, with data from the Framingham Offspring Study. The authors found that participants who had metabolic syndrome and exhibited subclinical disease had a risk of CVD that was 2.5-fold higher (HR, 2.67; 95% CI, 1.62-4.41) than that of those without metabolic syndrome or subclinical disease. The association of metabolic syndrome and CVD was attenuated in those without subclinical disease (HR, 1.59; 95% CI, 0.87-2.90).

380 found to predict future risk of myocardial infarction and stroke, and a change in carotid IMT has been validated as

I vascular marker for the progression of atherosclerosis. 81 Specifically, studies in patients with the metabolic syndrome I have demonstrated that carotid IMT abnormalities exist in patients with this syndrome and predict risk of CHD. Among 22,313 postmenopausal women, metabolic syndrome conferred an approximate threefold adjusted odds of subclinical carotid atherosclerosis, as measured by carotid IMT. 82 In these women, the metabolic syndrome but not BMI was associated with increased carotid IMT. Obesity had no independent effect, suggesting that metabolic abnormalities with metabolic syndrome aged 40 to 79 years developed carotid plague in a 5-year follow-up.

have also been shown to predict the progression of carotid in the ARIC study comparing those with versus those without twofold. metabolic syndrome, both prevalence of CHD (7.4% versus Computed Tomographic Angiography 3.6%) and average carotid IMT were significantly greater in those with versus those without metabolic syndrome. 86 Finally, even among nondiabetic young subjects from the Bogalusa Heart Study (n = 507), composite carotid IMT increased significantly with the number of metabolic syndrome components present, and metabolic syndrome predicted composite carotid IMT 75th percentile or higher by receiver operator characteristics curves. 87

Whereas the evidence relating metabolic syndrome to carotid IMT is strong, there remains debate as to whether carotid IMT can be used as a surrogate for assessment of the effects of therapy. Although some prevention trials with lipidlowering medications that used carotid IMT as a surrogate endpoint have shown that retardation in the progression of carotid IMT is accompanied by a reduction of clinical cardiovascular endpoints, 88,89 others have not shown this.

Coronary Artery Calcification

The presence and extent of coronary artery calcification (CAC) strongly correlate with the overall magnitude of coronary atherosclerosis plaque burden and with the development of subsequent coronary events. 90,91 The authors have previously demonstrated the presence of metabolic syndrome to be independently associated with an increased likelihood of CAC (compared with those without metabolic syndrome) and those with diabetes to have the highest likelihood of CAC. 92 Moreover, the prevalence of calcium among women with metabolic syndrome was as high as in those with diabetes. Metabolic syndrome without diabetes was independently associated with an increased likelihood of CAC. Similarly, the National Heart, Lung, and Blood Institute Family Heart Study has demonstrated metabolic syndrome to be independently associated with an increased likelihood of CAC and abdominal aortic calcification in both men and women after adjustment for other risk factors. 93

The Dallas Heart Study assessed the association between metabolic syndrome, diabetes mellitus, and subclinical atherosclerosis defined as CAC or abdominal aortic plaque detected by magnetic resonance imaging. Among 2735 participants, the prevalence of CAC was increased from those with neither metabolic syndrome nor diabetes (16.6%) to metabolic syndrome only (24%), to diabetes only (30.2%), to those with both metabolic syndrome and diabetes (44.7%). After adjustment, metabolic syndrome and diabetes were each independently associated with CAC. 94 Analysis of abdominal aortic plaque showed similar results, with the highest prevalence of subclinical atherosclerosis in those with both diabetes and metabolic syndrome.

In addition to having independent effects of other traditional mediate the risk of subclinical atherosclerosis. Prospective data risk factors, metabolic syndrome has a synergistic effect. A crossalso show increased carotid IMT in those with metabolic sectional study examining the combined effect of high LDL-C and syndrome. Bonera and associates 83 reported that 51% of people metabolic syndrome on CAC found that CAC in asymptomatic men with moderate or high LDL-C was magnified in persons with metabolic syndrome. 95 LDL-C was more strongly associated with An increasing number of metabolic syndrome risk factors subclinical atherosclerosis when subjects had metabolic syndrome.

Not only metabolic syndrome but high-normal fasting blood IMT in elderly women during a 12-year period. 84 In this glucose concentration has been shown to have increased levels of prospective study, the more metabolic syndrome risk factors subclinical atherosclerosis. In one study, high-normal fasting blood that developed during the 12-year period, the greater was the glucose concentration was found to be associated with increased increase in mean carotid IMT. Incident metabolic syndrome CAC in asymptomatic nondiabetic men. 96 The authors found that was a stronger predictor of subclinical atherosclerosis than high-normal fasting blood glucose concentration was associated were individual components of the syndrome. Metabolic - with CAC independent of metabolic syndrome. In another study, syndrome was associated with progression of carotid IMT these authors found an association of metabolic syndrome with even after the Framingham risk score was accounted for. CAC in asymptomatic men independent of the Framingham risk Finally, a cross-sectional analysis of 14,502 patients in the score. 97 Metabolic syndrome was present in 24% of the study -ARIC study demonstrated that the metabolic syndrome is participants. The prevalence of CAC increased with increasing associated with increased average carotid IMT. 85 In addition, number of metabolic syndrome risk factors. The presence of in a separate investigation of 14,502 black and white subjects metabolic syndrome increased the risk of any CAC by almost

Recent advances in contrast-enhanced computed tomo graphic angiography allow the direct visualization of calcified and noncalcified plaque. There are now some data showing that assessment of plaque by this method strongly correlates with cardiovascular events. 98 Using data from the ROMICAT study, Butler and colleagues recently showed that those with metabolic syndrome have a higher prevalence of coronary plaque than do those without metabolic syndrome. 99 The presence of any, calcified, and noncalcified plaque was higher in patients with than without metabolic syndrome (91%, 74%, and 77% versus 46%, 45%, and 40% of coronary segments with plaque, respectively). Metabolic syndrome was independently associated with both the presence and extent of overall plaque after adjustment for the Framingham risk score (odds ratio, 6.7). However, given the current radiation dose from computed tomographic angiography, routine use in asymptomatic patients should be avoided.

Ankle-Brachial Index and Peripheral Arterial Disease

A low ankle-brachial index has been previously shown to strongly predict morbidity and mortality in persons without known CVD. 100,101 There are limited data on the association of ankle-brachial index with metabolic syndrome. The study population of this crosssectional survey consisted of 502 patients recently diagnosed with CHD, 236 with stroke, 218 with peripheral arterial disease, and 89 with abdominal aortic aneurysm. The prevalence of the metabolic syndrome in the study population was 45%. In patients with peripheral arterial disease, this was 57%; in CHD patients, 40%; in stroke patients, 43%; and in patients with abdominal aortic aneurysm, 45%. Patients with the metabolic syndrome more often had a decreased ankle-brachial index (14% versus 10%; P = 0.06). 102

Myocardial Perfusion and Imaging of Inflammation

In addition to direct (multidetector computed tomographic angiography) and indirect (CAC) assessment of coronary plaque, coronary perfusion has also been shown to vary with metabolic syndrome. Some initial observations in this regard include the finding that among persons with metabolic abnormalities (metabolic syndrome or diabetes), there is an increased likelihood of myocardial ischemia, as assessed by nuclear single-photon emission computed tomography (SPECT), at intermediate levels of CAC (eg, 100 to 399), similar to that of those without such abnormalities who have higher levels of CAC (eg, > 400). 103 Also, the number of metabolic syndrome abnormalities increases the amount of ischemic area on stress SPECT. 104

In addition to perfusion imaging, we can now assess the degree of inflammation in different vascular beds. Fluorode oxyglucose (FDG) positron emission tomography can measure inflammation within the aorta and carotid arteries. Interscan plaque FDG variability during 2 weeks was very low, with high intraclass correlation (0.79-0.92) and high intraobserver agreement across most territories, suggesting its usefulness as a noninvasive plaque syndrome. Am J Cardiol 91:1421, 2003.) imaging technique for use in drug intervention studies. 105 A study showed that FDG uptake was significantly associated with waist circumference, HDL-C, HOMA insulin resistance, hsCRP, and a number of metabolic syndrome components (P < 0.05 to P < 0.001). waist circumference, blood pressure, and triglyceride levels and

Other Subclinical Disease Measures

Others have also examined the relationship of metabolic syndrome to other measures of subclinical atherosclerosis. In a random sample of 1153 French adults aged 35 to 65 years, the presence of metabolic syndrome was independently associated with the number of carotid and femoral plaques, carotid IMT, and pulse wave velocity, with approximately 80% of coronary artery disease events could odds ratios ranging from 1.8 to 2.15 by use of the NCEP definition potentially be prevented by control of blood pressure, LDL-C, and and 1.48 to 1.97 with the WHO definition. 107 A recently published investigation from the ARIC study demonstrated a stepwise without diabetes (Fig. 22-8). 112 gradient in echocardiographic left ventricular mass by increasing with metabolic syndrome, there were progressive increases in left benefit. ventricular mass and decreases in tissue Doppler imaging of diastolic function. 110 Others, however, have demonstrated that Lifestyle Modification there is greater impairment of global left ventricular function in patients with versus without metabolic syndrome based on an index of myocardial performance. 111

MANAGEMENT OF METABOLIC SYNDROME

The treatment goal for metabolic syndrome is to prevent future development of type 2 diabetes and diabetes-related cardiovascular first year, with continued weight loss to achieve a BMI < 25 kg/m² complications. Lifestyle modification is the mainstay therapy for , is noted. In addition, regular moderate-intensity physical activity metabolic syndrome. The NCEP ATP III included metabolic of at least 30 minutes and preferably 60 minutes a day at least 5 days syndrome within its lipid management guidelines to reinforce per week is recommended. Dietary recommendations focus in lifestyle therapies, including weight reduction, antiatherogenic diet, particular on reducing intakes of saturated fat, trans -fat, and and increased physical activity. The treatment approach focuses on cholesterol. glycemic control and control of cardiovascular risk factors, mainly with metabolic syndrome (exclusive of diabetes) have increased

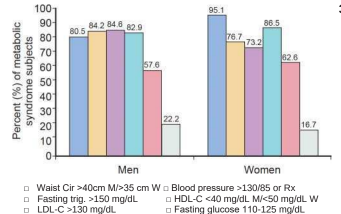


FIGURE 22-7 Prevalence of metabolic syndrome risk factors among US adults with metabolic syndrome (but without diabetes), National Health and Nutrition Examination Survey 1988-1994. (Modified from Wong ND, Pio JR, Franklin SS, et al: Preventing coronary events by optimal control of blood pressure and lipids in patients with the metabolic

depressed HDL-C concentration, 58% of men and 63% of women with metabolic syndrome also have levels of LDL-C of 130 mg/dL (3.4 mmol/L) or higher 112 (Fig. 22-7). The potential benefits from optimal control of lipids and blood pressure, in particular, can be significant. We have additionally shown (by statistically controlling lipids and blood pressure with Framingham risk algorithms) that HDL-C to optimal levels in persons with the metabolic syndrome

Mounting evidence suggests that lifestyle modification with number of metabolic syndrome disorders (none, any, two, or all weight loss and increased physical activity is beneficial, although three risk factors) in both men and women. 108 Moreover, among specific studies in metabolic syndrome are needed. There are overweight hypertensive patients, those with versus without suggestions from the Finnish Diabetes Prevention Study that metabolic syndrome had significantly greater left ventricular mass individuals with metabolic syndrome show less development of even after controlling for age, gender, and blood pressure. 109 diabetes with lifestyle advice. 113 In many people, however, Finally, in a study of 607 adults with normal left ventricular function pharmacological intervention will be needed. Long-term studies assessed by echocardiogra phy, whereas left ventricular ejection will help establish whether existing or newer agents, such as fraction was similar among normals, those with one or two agonists for the peroxisome proliferator-activated (PPAR) a and y metabolic syndrome criteria (pre-metabolic syndrome), and those receptors or cannabinoid 1 receptor blockers, could be of specific

Intensive lifestyle modification is the mainstay of treatment in lowrisk patients. The most recent (2005) American Heart Association/National Heart, Lung, and Blood Institute statement 20 on the diagnosis and management of metabolic syndrome noted specific lifestyle-related therapeutic targets to include abdominal obesity, physical inactivity, and diet (Table 22-2). For abdominal obesity, a goal of reducing body weight by 7% to 10% during the

No single diet is currently recommended for patients with the hypertension and hyperlipidemia. Although most persons (>75%) metabolic syndrome, although evidence suggests a lower prevalence of metabolic syndrome with dietary patterns rich in fruit, vegetables, whole grains, dairy products,

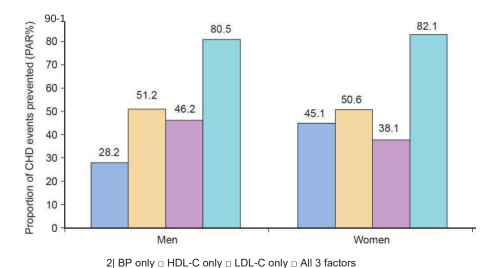


FIGURE 22-8 Proportion of CHD events potentially preventable from optimal control of blood pressure (to <120 mm Hg/<80 mm Hg), LDL-C (to <100 mg/dL), and HDL-C (to >60 mg/dL) in men and women. (Modified from Wong ND,

Pio JR, Franklin SS, et al: Preventing coronary events by optimal control of blood pressure and lipids in patients with the metabolic syndrome. Am J Cardiol 91:1421, 2003.)

and unsaturated fats. The literature favors diets that are lower in Pharmacological Treatment carbohydrates and higher in healthy unsaturated fats and protein, compared with traditional high-carbohydrate and low-fat diets, to treat obesity, to reduce triglycerides, and to increase HDL-C. Diets that foster adherence, such as those modeled after the Mediterranean diet, not only improve risk factors but also reduce coronary mortality after myocardial infarction. 114 For example, recent evidence indicates that a Mediterranean-style diet reduces the prevalence of the metabolic syndrome while improving endothelial function. 115 Moreover, a Mediterranean diet is associated with increasing longevity 116 and improved survival after proinflammatory states (Table 22-3). myocardial infarction. 117

In a study using data from NHANES III, participants diagnosed with metabolic syndrome had a lower consumption of fruit, vegetables, and antioxidants than did those without metabolic syndrome. 118 The Coronary Artery Risk Development in Young Adults (CARDIA) study showed that consumption of dairy products was associated with a significantly reduced risk of metabolic syndrome. 119 The Framingham Off spring Study showed that whole-grain and cereal fiber intakes were associated with reduced risk of metabolic syndrome. 120 Importantly, both the US Diabetes Prevention Program 121 and the Finnish Diabetes Prevention Study 122 showed a 58% lower risk for the development of new onset diabetes in pre diabetic individuals resulting from prescribed weight reduction and physical activity.

Lifestyle and behavior modification can be difficult. O'Malley and coworkers 123 examined the effect of case man agement on the development of the metabolic syndrome. In a randomized control trial with 450 patients, they were able to demonstrate greater that those with metabolic syndrome derive a disproportionately improvement in motivation to change behavior and lower prevalence of metabolic syndrome in 6 months after intervention with a cardiovascular case management program.

In severely obese patients in whom lifestyle measures are not sufficient to adequately address the problem, pharmaco logic therapy for weight loss (see Chapter 19) can be recommended as an adjunct. Finally, laparoscopic weight reduction surgery for the sole purpose of treating metabolic syndrome, although not recommended, has been shown in studies that had follow-up after this procedure to result in reversal of metabolic syndrome in 80% 124 to 96% 125 of patients.

When lifestyle modification fails and in high-risk patients, medications that target individual risk factors are recommended -(eg, antihypertensives, lipid-lowering drugs, hypo glycemic drugs, antiplatelet drugs, and weight loss drugs). Specific therapeutic targets recommended by the American Heart Association/National Heart, Lung, and Blood Institute statement 20 on metabolic syndrome focus on atherogenic dyslipidemia, elevated blood pressure, elevated glucose, and prothrombotic

Dyslipidemia

For atherogenic dyslipidemia, LDL-C remains the primary target of therapy. Goals are consistent with the NCEP guide lines, namely, < 100 mg/dL in high-risk patients (with an option for < 70 mg/dL), <130 mg/dL in moderately high risk subjects (with an option for < 100 mg/dL), < 130 mg/dL in moderate-risk patients, and < 160 mg/dL in lower-risk patients. In addition, non-HDL-C is a secondary target for therapy when triglycerides are 200 mg/dL or greater; goals are 30 mg/dL higher than the respective LDL-C targets. HDL-C remains a tertiary target of therapy; lifestyle or pharmacological therapy is recommended to raise HDL-C levels < 40 mg/dL in men or < 50 mg/dL in women (see Table 22-3).

Treatment of dyslipidemia associated with metabolic syndrome with fibrates or niacin, especially in combination with statins, has been shown to be effective in addressing elevated triglycerides and low HDL-C, which are highly prevalent in persons with metabolic syndrome. Post hoc analysis of several of the fibrate studies shows large reduction in cardiovascular events with treatment by these agents. 126 A meta analysis of the Familial Atherosclerosis Treatment Study (FATS), the HDL-Atherosclerosis Treatment Study (HATS), and the Armed Forces Regression Study (AFREGS) showed that patients with metabolic syndrome had 50% more rapid coronary stenosis progression than did those without metabolic syndrome and that combination of lowering LDL-C plus increasing HDL-C resulted in added benefit. 127 Combination therapy with a statin and fibrate or niacin resulted in a 54% decrease in cardiovascular events in those with metabolic

TABLE 22—2 Treatment of Lifestyle Risk Factors for Long-Term Prevention of Atherosclerotic Cardiovascular Disease or Prevention and Treatment of Type 2 Diabetes

Therapeutic Target and Goals of Therapy Therapeutic Recommendations

Abdominal Obesity

Goal: Reduce body weight by 7%-10% during first year of therapy. Continue weight loss thereafter to extent possible, with goal to ultimately achieve desirable weight (BMI <25 kg/m²)

Consistently encourage weight maintenance or reduction through appropriate balance of physical activity, calorie intake, and formal behavioral programs when indicated to maintain or to achieve waist circumference of <40 inches in men and <35 inches in women. Aim initially at slow reduction of 7% to 10% from baseline weight. Even small amounts of weight loss are associated with significant health benefits.

Physical Inactivity

Goal: Regular moderate-intensity physical activity; at least 30 min of continuous/intermittent (preferably 60 min) 5 days/wk, but preferably daily

In patients with established CVD, assess risk with detailed physical activity history or exercise test to guide prescription. Encourage 30-60 min moderate-intensity aerobic activity (e.g., brisk walking), preferably daily, supplemented by increase in daily lifestyle activities (e.g., pedometer step tracking, walking breaks at work, gardening, household work). Higher exercise times can be achieved by accumulating exercise throughout day. Encourage resistance training 2 days/wk. Advise medically supervised programs for high-risk patients (e.g., recent acute coronary syndrome or revascularization, congestive heart failure).

Atherogenic Diet

Goal: Reduced intakes of saturated fat, transfat, cholesterol

Recommendations: Saturated fat <7% of total calories: reduce trans-fat: dietary cholesterol <200 mg/ day; total fat 25%-35% of total calories. Most dietary fat should be unsaturated; simple sugars should be limited.

Modified from Grundy SM: Metabolic syndrome scientific statement by the American Heart Association and the National Heart, Lung, and Blood Institute. Arterioscler Thromb Vasc Biol 25:2243, 2005.

interaction of statins with gemfibrozil.

Elevated Glucose Concentration

The goal is to delay progression to type 2 diabetes for patients with impaired fasting glucose or to reduce the HbA1c to < 7.0% if the patient is diabetic (although recent guidelines have suggested less stringent guidelines for those with long standing diabetes or diabetes complicated by other comor bidities including CVD). Weight reduction and increased physical activity remain the primary intervention for persons with impaired fasting glucose, and in those with diabetes, pharmacotherapy should be supplemented as needed to reach HbA1c goals (see Table 22-3).

control are similar to those for patients with diabetes.

and PPAR y agonists (thiazolidinediones).

In persons with diabetes, fibrates and thiazolidinediones have

syndrome each 10% decrease in LDL-C or 10% increase in HDL-C also been shown to have benefits. In the recent PROAC TIVE clinical was associated with an 11% or 22% event risk reduction, trial, although the primary composite endpoint was not respectively. The evidence supports treatment of dyslipidemia significantly reduced in those receiving pioglitazone, the main associated with metabolic syndrome, beyond statin therapy for secondary endpoint of myocardial infarction, stroke, or CVD death LDL-C, to include fibrates or niacin, used in combination with was significantly reduced in the group receiving pioglitazone. 134 statins to incrementally improve HDL-C and triglycerides in Also, in the Fenofibrate Intervention and Event Lowering in addition to LDL-C. The safety of combination lipid therapy is better Diabetes (FIELD) study, although the effect of fenofibrate on the established, with fenofibrate lacking the well-known strong primary study endpoint of CHD events in the entire trial was not significant, there was a reduction in CVD events in those with low HDL-C, high tri glycerides, or hypertension. 135 The thiazolidinediones lessen insulin resistance and modestly improve various metabolic risk factors. Moreover, the thiazolidinedione rosiglitazone in the Diabetes Reduction Assessment with Ramipril Rosiglitazone Medication (DREAM) trial and the thiazolidinedione pioglitazone in the ACT-NOW study, administered to subjects with prediabetes, showed significant reductions in the onset of diabetes. 136,137

Elevated Blood Pressure

Blood pressure goals call for a reduction of blood pressure to at least Metformin, a biguanide, improves insulin sensitivity and < 140/90 mm Hg (or < 130/80 mm Hg if diabetes is present); and if decreases hepatic glucose output. The Diabetes Prevention Program blood pressure is at or above 120/80 mm Hg, lifestyle modification demonstrated that metformin is effective at slowing the onset of is initiated or maintained by weight control, increased physical diabetes in those with impaired glucose tolerance, but participants activity, alcohol moderation, and sodium restriction. Pharmacologic taking metformin in the Diabetes Prevention Program trial did not therapy should be supple mented as needed when blood pressure have significant resolution of metabolic syndrome compared with is > 140/90 mm Hg (see Table 22-3), although more recent those taking placebo. 128 Studies done with rimonabant (which was guidelines call for lower targets of < 130/80 mm Hg in persons at marketed for only a brief period in Europe and never came to intermediate or higher risk with multiple risk factors whose 10-year market in the United States) did show approximately one third of risk of CHD exceeds 10% (many metabolic syndrome patients fit participants taking the drug for 1 year to have resolution of this criterion). 138 A recently recognized effect of angiotensin metabolic syndrome. 129-131 The goals for cardiovascular risk factor converting enzyme inhibitors and angiotensin receptor blockers is a consistent reduction in the incidence of new-onset diabetes among Currently, only metformin is recommended as an option for patients with essential hypertension. Thus, pharmacological therapy in persons with impaired fasting glucose; there are no such blockade of the RAS, in addition to having proven benefits in recommendations for thiazolidinediones or other antidiabetic reducing cardiovascular events and reducing the progression of agents at present. 132,133 Drugs that could potentially target insulin renal disease, may also be able to prevent the development of resistance include weight loss drugs, PPAR a agonists (fibrates), diabetes. Elucidation of the mechanisms by which these drugs prevent or delay diabetes might open the door to new therapeutic strategies.

Therapeutic Targets and Goals of Therapy

Atherogenic Dyslipidemia

Primary target: LDL-C

Reduces LDL-C levels to ATP III goals (see Therapeutic Recommendations).

Secondary target: non-HDL-C

for atherogenic dyslipidemia.

If TG > 200 mg/dL, reduce non-HDL-C to ATP III goals (after attaining LDL-C goals; see Therapeutic Recommendations). Tertiary target: HDL-C If HDL-C < 40 mg/dL in men or < 50 mg/dL in women after reaching non-HDL-C goal, raise HDL-C to the extent possible with standard therapies

Therapeutic Recommendations

For elevated LDL-C: Give priority to reduction of LDL-C over other lipid parameters. Achieve LDL-C goals based on patient's risk category.

LDL-C goals for different risk categories:

High risk*: < 100 mg/dL (optional < 70 mg/dL for very-high-risk patients \cdot) Moderately high risk \cdot : < 130 mg/dL (optional < 100 mg/dL) Moderate risk $_{\$}$: < 130 mg/dL Lower risk $_{\$}$: < 160 mg/dL

If TG > 200 mg/dL, goal for non-HDL-C for each risk category is 30 mg/dL higher than for LDL-C. If TG > 200 mg/dL after achieving LDL-C goal, consider additional therapies to reach non-HDL-C goal.

For reduced HDL-C: If HDL-C is low after achieving non-HDL-C, either lifestyle therapy can be intensified or drug therapy can be used for raising HDL-C levels, depending on the patient's risk category.

Elevated BP

Reduce BP to at least achieve BP of < 140/90 mm Hg (or < 130/80 mm Hg if diabetes is present). Reduce BP further to the extent possible through lifestyle changes.

For BP > 120/80 mm Hg: Initiate or maintain lifestyle modification by weight control, increased physical activity, alcohol moderation, sodium reduction, and emphasis on increased consumption of fresh fruits, vegetables, and low-fat dairy products in all patients with metabolic syndrome. For BP > 140/90 mm Hg (or > 130/80 mm Hg if diabetes is present), add BP medication as needed to achieve goal BP.

Elevated Glucose

For IFG, delay progression to type 2 diabetes mellitus. For diabetes, hemoglobin A1c < 7%.

For IFG, encourage weight reduction and increased physical activity. For type 2 diabetes, lifestyle therapy and pharmacotherapy, if necessary, should be used to achieve near-normal HbA1c (< 7%). Modify other risk factors and behaviors (eg, abdominal obesity, physical inactivity, elevated BP, lipid abnormalities).

Prothrombotic State

Reduces thrombotic and fibrinolytic risk factors

Proinflammatory states

For high-risk patients, initiate and continue low-dose aspirin therapy; in patients with ASCVD, consider clopidogrel if aspirin is contraindicated. For moderately high risk patients, consider low-dose aspirin prophylaxis.

Recommendations: No specific therapies beyond lifestyle therapies

Two large clinical trials, the Heart Outcomes Prevention Evaluation (HOPE) study (ramipril) and Losartan Intervention For Endpoint (LIFE) reduction in hypertension study (losartan), have demonstrated the benefit of blocking the RAS with either an angiotensin-converting enzyme inhibitor or angiotensin receptor blocker and the improvement in insulin sensitivity. ^{139,140} Other clinical studies have shown that inhibition of the RAS with either angiotensin-converting enzyme inhibitors or angiotensin receptor blockers results in increasing levels of adiponectin, which is associated with improved insulin sensitivity. ¹⁴¹

A prospective long-term randomized trial assessed the effectiveness of an angiotensin receptor blocker in reducing the incidence of diabetes among those with metabolic syndrome . Results from the DREAM trial concluded that among persons with impaired fasting glucose levels or impaired glucose tolerance, the use of ramipril (compared with placebo) did not significantly reduce the incidence of diabetes or death but did improve normoglycemia. 136

Prothrombotic and Proinflammatory States

To reduce thrombotic risk, low-dose aspirin therapy is initiated or continued in moderately high risk (eg, when 10-year risk of CHD is 10% or greater) or high-risk patients; and in those with CVD, clopidogrel is considered if aspirin is contraindicated. For proinflammatory states, there are no specific recommended therapies beyond lifestyle modifications, however (see Table 22-2).

CONCLUSION

The metabolic syndrome is associated with an increased risk for the development of diabetes, CHD, and CVD. Whereas diabetes is frequently regarded as a CHD risk equivalent, a wide spectrum of risk for CHD is present in persons with metabolic syndrome (but without diabetes), necessitating careful assessment of cardiovascular risk. Although initial global risk assessment can use Framingham or other risk algorithms,

consideration should be given to stratification of risk by the presence and extent of other metabolic syndrome risk factors, novel risk markers, and measures of subclinical atherosclerosis.

Health care providers should regularly assess for the presence of multiple risk factors as well as metabolic syndrome at regular intervals in each patient. Identification of the metabolic syndrome as a clinical condition will also accustom physicians to simultaneous treatment of multiple risk factors (particularly abdominal obesity, dyslipidemia, and elevated blood pressure), instead of the traditional model of treatment of risk factors in isolation. Most importantly, intensified efforts at lifestyle therapies, including effective dietary and physical activity counseling by trained individuals who can provide the necessary guidance and follow-up, are needed and crucial if significant strides are to be made in reducing the prevalence of metabolic syndrome and its future diabetic and cardiovascular complications.

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SECTION VII

Special Populations

CHAPTER 23

Role of Ethnicity in Cardiovascular Disease: Lessons Learned from MESA and Other Population-Based Studies

KEY POINTS
Karol E. Watson and Ashkan Afshin

- Cardiovascular morbidity and mortality rates differ among diverse racial and ethnic groups in the United States.
- The major cardiac risk factors are the same in all ethnic and racial groups in the United States and worldwide; however, the demographics and relative weight attributed to each factor may differ.
- Chronic, subclinical inflammation appears to be one pathophysiologic mechanism explaining the increased risk of atherosclerotic disease regardless of the amount of obstruction produced by that coronary disease, and African Americans are more likely than white Americans to carry allelic variants demonstrated to increase production of inflammatory cytokines.
- Framingham risk calculation performs reasonably well for prediction of CHD events in all racial and ethnic groups; however, among Japanese American and Hispanic men and Native American women, the Framingham functions systematically overestimated the risk of 5-year CHD events.
- Different races respond differently to a variety of cardiovascular medications in terms of both efficacy and toxicity.

CARDIOVASCULAR DISEASES IN RACIAL AND ETHNIC MINORITIES: OVERVIEW AND PERSPECTIVES

Cardiovascular disease (CVD) is the leading cause of death in the United States and worldwide. ¹ In 2001, heart disease accounted for approximately 29% of deaths among US residents; 17% of those deaths occurred among persons aged < 65 years. ² From 1996 to 2006, death rates from CVD have decreased 29%; however, the decline has not been uniform for all populations. ³ It is well documented that cardiovascular morbidity and mortality rates differ among various racial and ethnic groups in the United States. ⁴

Recent data from the Centers for Disease Control and Prevention reveal that African Americans have earlier and higher mortality rates from coronary heart disease (CHD) than those of whites, American Indian/Alaska Natives, Asian/Pacific Islanders, or Hispanics ⁴ (Fig. 23-1). Ethnic differences in the prevalence of complex diseases such as atherosclerosis are undoubtedly multifactorial . It is clear that there are social, environmental , biological, genetic, and probably other determinants leading to the disappearance of CHD.

Whereas earlier articles have explored many of these factors, ongoing large population -based and cohort studies are lending valuable insight. One of these studies, the Multi-Ethnic Study of Atherosclerosis (MESA), is adding important insights into

CVD among various ethnic groups in the United States. MESA was initiated in July 2000 to investigate the prevalence, correlates, and progression of subclinical CVD in a population-based sample of 6814 men and women aged 45 to 84 years. The cohort was selected from six United States field centers and is approximately 38% white, 28% African American, 23% Hispanic, and 11% Asian (primarily of Chinese descent). Base line measurements taken included measurement coronary calcium by computed tomography, measurement of ventricular mass and function by cardiac magnetic resonance imaging, measurement of flowbrachial artery endothelial vasodilation, carotid intima-media wall thickness, and measurement of peripheral vascular disease by ankle and brachial blood pressures. Assessments of demographic measures, standard CVD risk factors, sociodemographic factors, life habits, and psychosocial factors were also made. Blood samples were assayed for blood chemistries, lipids, inflammatory markers, and DNA.

The MESA cohort has been observed since 2000 for identification and characterization of CVD events, including acute myocardial infarction and other CHD, stroke, peripheral arterial disease, heart failure, therapeutic interventions for CVD, and mortality. Thus, MESA and other epidemiological studies will lend invaluable insights into the role of race and ethnicity in CVD.

This chapter summarizes currently - available data on ethnic differences in CVD,

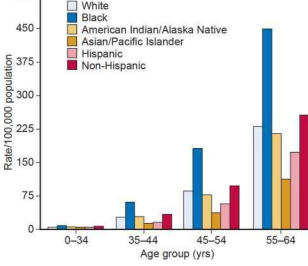


FIGURE 23-1 The 2001 CHD mortality rates among persons <65 years from different racial and ethnic groups. African Americans (blue bar) have higher CHD rates compared with all other groups. (From Centers for Disease Control and Prevention [CDC]: Disparities in premature deaths from heart disease—50 States and the District of Columbia, 2001. MMWR Morb Mortal Wkly Rep 53:121, 2004.)

cardiac risk factors, vascular biology, genetic factors, and socioeconomic determinants of atherosclerosis, but it is important to keep in mind that ethnic populations are far from homogeneous with respect to many genetic and biological traits. ⁵ Nevertheless, investigation into variations among ethnic groups can be an important key to understanding the causes of complex genetic conditions like atherosclerosis. Such investigations can also lead to important research into pathophysiology and treatment, thus improving outcomes for all populations.

EPIDEMIOLOGY OF CARDIOVASCULAR RISK FACTORS AMONG RACIAL AND ETHNIC POPULATIONS IN THE UNITED STATES

Considerable information has been gathered cardiovascular risk factors in ethnic populations. The major cardiac risk factors are the same in all ethnic and racial groups in the United States and worldwide; however, the demographics and relative weight attributed to each factor may differ. Among African Americans, for example, some of the excess CHD may be accounted for by the excess prevalence of known complicating risk factors, such as hypertension, left ventricular hypertrophy, and diabetes mellitus. Differences in other risk factors, such as plasma lipoprotein profiles, may not completely explain disparities but may lend insight into ethnic differences in the biology of atherosclerosis.

Hypertension

Worldwide, hypertension is known to account for considerable cardiovascular risk. ⁶ The INTERHEART trial highlighted this risk. ⁷ In the INTERHEART study, a standardized case control study that screened all patients admitted to the coronary care unit or equivalent cardiology ward for a first myocardial infarction at 262 participating centers in 52 countries throughout Africa, Asia, Australia, Europe, the Middle East, and North and South America, hypertension was thought to account for 23.4% of the population-attributed risk for a myocardial infarction.

Compared with all other ethnic groups in the United States,

hypertension in African Americans is more common, begins earlier, is more severe, and causes more target organ damage. $^{8-12}$ Relative to hypertensive whites, African Americans with hypertension demonstrate delayed sodium excretion , plasma volume expansion, lower plasma renin activity, elevated intracellular sodium concentration, and altered numbers and activities of sodium transporters (Na \cdot -H \cdot anti-t $_{\rm NT}$ \cdot K $^{+}$ - ATP d $_{\rm NT}$ -KN Cl $_{\rm T}$ tt) 13 , 14 ponies, Na ,K -AiPase, an Na -K -C conanspoiieij. Many of these

differences may have a genetic basis.

To possibly explain the greater prevalence and severity of hypertension in African Americans, several candidate genes have been explored. Genes encoding components of epithelial sodium channels, ¹⁵ the renin-angiotensin-aldosterone system, ¹⁶⁻¹⁸ alpha- and beta-adrenergic receptors, ^{19,20} endothelin, ^{21,22} kallikrein, ²³ natriuretic peptides, ²⁴ and the nitric oxide pathway ²⁵ have been investigated. To date, few convincing genetic explanations for ethnic differences have emerged.

Hypertension in part increases CHD risk by predisposing to left ventricular hypertrophy. ²⁶ Left ventricular hypertrophy increases cardiac risk up to fourfold and is more common in African Americans, even after adjustment for blood pressure . The mechanisms by which left ventricular hypertrophy increases risk are poorly understood but likely include predisposition to

ischemia and arrhythmias. 26 The MESA has given us valuable insight into the prevalence, treatment, and control of hypertension in the United States. Whereas most prior studies investigating the association between ethnicity and hypertension in the United States focused on differences between African Americans and whites and did not include other racial and ethnic groups such as Chinese or Hispanics, a paper by Kramer and colleagues ²⁷ used MESA data to examine the association between ethnicity and hypertension and hypertension treatment among white, African American, Chinese, and Hispanic ethnic groups. These authors found that the prevalence of hypertension, defined as systolic blood pressure < 140 mm Hg and/or diastolic blood pressure < 90 mm Hg or self-reported treatment for hypertension, was significantly higher in African Americans compared with whites (60% versus 38%; P < 0.0001), whereas prevalence in Hispanic (42%) and Chinese participants (39%) did not differ significantly from that in whites. After adjustment for age, body mass index (BMI), prevalence of diabetes mellitus, and smoking, African American ethnicity (OR, 2.21; 95% CI, 1.91-2.56) and Chinese ethnicity (OR, 1.30; 95% CI, 1.07-1.56) were significantly associated with hypertension compared with whites. They further found that among hypertensive MESA participants, the percentage of treated but uncontrolled hypertension in whites (24%) was significantly lower than that in African Americans (35%; P <0.0001), Chinese (33%; P = 0.003), and Hispanics (32%; P =0.0005), but only African American race and ethnicity remained significantly associated with treated but uncontrolled hypertension after controlling for socioeconomic factors (OR, 1.35; 95% CI, 1.07-1.71).

Dyslipidemia

The epidemiological relationship between serum cholesterol levels and the risk of CHD is well documented. ²⁸⁻³² Across cultures, cholesterol is linearly related to CHD mortality, and despite differences in the prevalence of CHD between various populations, the relative increase in CHD mortality rates for a given cholesterol increase is remarkably consistent. ³² There is a complex interplay between genetic and environmental factors that influence the expression of lipoprotein levels in individuals and between groups of people, and differences in plasma lipoprotein levels have been reported between various ethnic groups.

African Americans have one of the highest CHD event rates of any ethnic or racial group in the United States. 33,34 Despite this fact, epidemiological studies have consistently shown that plasma lipoprotein concentrations appear more favorable in African Americans than in white Americans. 35-38

African Americans have been shown to have higher highdensity lipoprotein cholesterol (HDL-C) levels than white populations. This is likely due, at least in part, to the lower activity of hepatic lipase in African Americans. ³⁹ Hepatic lipase is an enzyme involved in HDL-C catabolism; thus, the lower the hepatic lipase activity, the higher the HDL-C level. Hepatic lipase activity has been found to be lower in African American men than in white men by Vega and colleagues, ³⁹ in part because of increased prevalence of a hepatic lipase allele (514T) that is associated with reduced hepatic lipase activity.

Two other polymorphisms that cause amino acid substitutions in hepatic lipase (N193S and L334F) have since been found that are also associated with lower hepatic lipase activity and are also much more common in African Americans than in whites. 40 This higher HDL-C level, however, may not protect African Americans from CHD, as one might expect. In an analysis of the Veterans Affairs HDL Intervention Trial (VA-HIT), Rubins and colleagues 41 found that the African American participants, all of whom had CHD, had low-density lipoprotein cholesterol (LDL-C) levels similar to those of the white men with CHD in the study but substantially higher HDL-C levels. Furthermore, compared with whites, African Americans have been found to have similar or slightly lower total cholesterol levels, lower LDL-C levels, and lower triglyc eride levels, 42,43 thus giving African Americans what would appear to be a more favorable lipoprotein profile.

Small, dense LDL-C particles are associated with an atherogenic lipoprotein phenotype that increases the risk of myocardial infarction up to three times. 44 In a study by Kral and colleagues, 45 racial differences in the prevalence of small, dense LDL-C were studied. They investigated 159 African American and 477 white siblings of persons with premature CHD (CHD occurring at < 60 years). Multiple logistic regression analysis demonstrated that white race (P = 0.009), tri glyceride level (P = 0.009) 0.0001), and diabetes (P = 0.02) were independent predictors of the likelihood of having small, dense LDL-C particles. White individuals had more small, dense LDL-C particles than African Americans did despite comparable levels of total cholesterol and

Lipoprotein(a) is structurally similar to LDL-C, with an additional disulfide-linked glycoprotein termed apolipoprotein(a). 46 Apolipoprotein(a) shares extensive structural homology with plasminogen but varies in size, which is due to the variation in the number of kringle 4-like domains (type 2 repeats) of plasminogen. Because of the size heterogeneity, apolipoprotein(a) exhibits a genetic size polymorphism with apparent molecular masses of isoforms ranging from 300 to 800 kDa. 47,48

There are considerable differences in the mean of plasma lipoprotein(a) concentrations between different populations and ethnic groups. 49 Many although not all epidemiological and case-control studies have shown that when lipoprotein(a) is present in high level in the plasma, it is an independent risk factor for CHD. 50 52 In addition to high lipoprotein(a) levels, the presence of small apolipoprotein(a) isoforms (with fewer kringle 4 type 2 repeats) has been associated with CHD in whites . 53,54 Interestingly, although mean lipoprotein(a) levels are more than twice as high in African Americans as in whites, some studies have failed to establish a significant association between elevated lipoprotein(a) levels and CHD among African Americans. 55,56 The reason for this may be that the majority of whites with high lipoprotein(a) levels possess at least one small apolipoprotein(a) isoform; however, the majority of African Americans with high lipoprotein(a) levels have no small apolipoprotein(a) isoforms.

The major Hispanic subgroups are Mexican Americans, Central and South Americans, Puerto Ricans, and Cuban Americans, with Mexican Americans making up the largest single Hispanic group. 59 Native Mexicans have lipids characterized by low HDL-C and elevated triglyceride levels. 60 In a survey done in 417 Mexican cities, information on lipid levels was obtained for 15,607 subjects, 20 to 69 years of age. Mean total cholesterol concentration in this cohort was 185 mg/dL; mean triglyceride level was 212 mg/dL; mean HDL-C level was 40 mg/dL; and mean LDL-C concentration was 118 mg/dL. The most prevalent lipoprotein abnormality in this cohort was low 23 HDL-C (HDL-C levels below 35 mg/dL), which occurred in 46% of men and 29% of women.

Hypertriglyceridemia (triglyceride levels > 200 mg/dL) was the second most prevalent abnormality, occurring in 24.3% of O. participants, with severe hypertriglyceridemia (> 1000 mg/ dL) m being observed in 0.42% of this population. Half of the subjects swith hypertriglyceridemia had a mixed dyslipidemia with low HDL-C levels as well. Insulin resistance was found to be prevalent in this population, being found in 59% of the subjects. 51 Thus, the prevalence of dyslipidemia in urban Mexican adults is very high, with much of this likely due to insulin resistance.

This pattern of dyslipidemia has also been seen in Mexican American adults and children living in the United States. 61 When Hispanics of different ancestry are studied, similar results are seen as well. Bermudez and colleagues 62 studied 490 Hispanics of Caribbean origin (Puerto Rico and the Dominican Republic) and 163 non-Hispanic whites. They found that concentrations of total cholesterol, HDL-C, and apolipoprotein AI were significantly lower among Hispanic women than among non-Hispanic white women. Although LDL-C concentrations are not higher in Hispanics, there appears to be a higher incidence of small, dense LDL-C.

In the Insulin Resistance Atherosclerosis Study, Hispanics not only had lower HDL-C and higher triglyceride levels than non-Hispanic whites did but also a smaller LDL-C particle size. 63 In regard to lipoprotein(a), available data suggest somewhat higher values in Hispanics compared with non-Hispanic whites. In a study by Chiu and associates, 64,390 non-Hispanic whites and 214 Hispanics from San Luis Valley, Colorado, were studied. Mean (\pm SD) and median lipoprotein(a) levels were 9.6 \pm 12.5 mg/dL and 3.8 mg/dL, respectively, in non-Hispanic whites and 12.1 \pm CL 15.6 mg/dL and 4.9 mg/dL, respectively, in Hispanics.

Asian Indians are known to be at increased risk for CHD. 65 zr Rates of coronary artery disease in young Asian Indians younger than 40 years are 3 to 10 times higher than those in other populations, and some of this increased risk may be conferred by C__ dyslipidemia. The typical lipid profile of Asian Indians living in Western societies is characterized by hyper triglyceridemia, low levels of HDL-C, and high levels of small, dense LDL-C. 66 In a study by Hoogeveen and colleagues, 67 Asian Indian subjects living in India and Asian Indians living in the United States were examined. Asian Indians living in the United States had higher plasma levels of triglyceride, total cholesterol, and LDL-C as well as lower HDL-C levels than did Asian Indians in India. Much of the dyslipidemia in Asian Indians may at least in part be due to the greater prevalence of insulin resistance observed in this population. 68

Small, dense LDL-C is also found more commonly in Asian Indians than in whites. In a study of 78 subjects, the prevalence of small, dense LDL-C was significantly higher in Asian Indians compared with white subjects (44% versus 21%; P < 0.05). 69 In this study, the increased prevalence of small, dense LDL-C type appeared to be due to the increased triglycerides.





</, similar values; ft, higher values; U, lower values

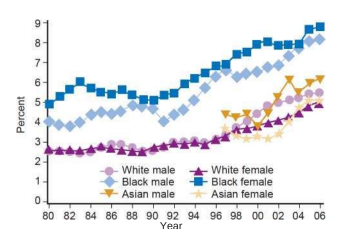


FIGURE 23-2 Age-adjusted percentage of civilian, noninstitutionalized population with diagnosed diabetes, by race and sex, United States, 1980-2006. Data show that blacks are disproportionately affected by diabetes. From 1980 through 2006, the age-adjusted prevalence of diagnosed diabetes increased among all sex-race groups examined. From 1980 through 2006, the age-adjusted prevalence of diagnosed diabetes was higher among blacks than whites and highest among black females. During this period, age-adjusted prevalence increased 116% among white males, 81% among white females, 100% among black males, and 69% among black females. Among Asians, the age-adjusted prevalence increased 42% among males and 38% among females from 1997 through 2006. (Data source: Centers for Disease Control and Prevention, National Center for Health Statistics, Division of Health Interview Statistics, data from the National Health Interview Survey. US Bureau of the Census, census of the population and population estimates. Data computed by the Division of Diabetes Translation, National Center for Chronic Disease Prevention and Health Promotion, Centers for Disease Control and Prevention.)

Higher serum lipoprotein(a) concentrations have also been reported in Asian Indians. In a study of young Asian Indian patients (younger than 45 years) who had suffered a myocardial infarction, the mean lipoprotein(a) level was 22.3 ± 5.4 mg/dL in patients and 9.3 ± 22.6 mg/dL in controls. 67.70

These data on ethnic differences in lipoprotein levels are summarized in Table 23-1. Of course, these general trends are based on population studies, and lipoprotein levels in a given individual may depart significantly. Nevertheless, knowledge of ethnic trends in plasma lipoprotein levels is useful for prognosis and treatment of CHD.

Insulin Resistance and Diabetes Mellitus

Type 2 diabetes varies considerably more by race and ethnicity as shown in Figure 23-2. The highest rates in the United States are seen in African Americans, Hispanic Americans, American Indians, and Asian/Pacific Islanders.

In the Atherosclerosis Risk in Communities (ARIC) study, the incidence of diabetes was 2.4-fold greater in African American women and 1.5-fold greater in African American men than in their white counterparts. ⁷¹ Excess adipose tissue accounted for almost half of the increased risk in African American women but little of the excess risk in African American men. African Americans with diabetes also have an increased risk of target organ damage, ^{72,73} and several studies have documented a

higher prevalence of insulin resistance in African Americans, even after correction for obesity and lifestyle factors. ⁷⁴

In 2000, of the 30 million Hispanic Americans, about 2 million had been diagnosed with diabetes. ⁷⁵ Hispanic Americans are about twice as likely to have diabetes as are non-Hispanic whites of similar age. Diabetes is particularly common among middle-aged and older Hispanic Americans. For those aged 50 years or older, about 25% to 30% have either diagnosed or undiagnosed diabetes. ⁷⁵ Diabetes is twice as common in Mexican American and Puerto Rican adults as in non-Hispanic whites. The prevalence of diabetes in Cuban Americans is lower, but it is still higher than that of non-Hispanic whites. ⁷⁵

Asian Americans have a diverse population composed of individuals of Chinese, Filipino, Asian Indian, Vietnamese, Korean, and Japanese descent and others with Asian ancestry, such as Pacific Islanders. ⁷⁶ Studies suggest that Asians are more likely to develop diabetes than are whites, even when they are less obese. For example, in the Honolulu Heart Study, the prevalence of physician-diagnosed plus newly diagnosed diabetes was 40% in Japanese American men older than 70 years ⁷⁷ compared with 19.2% in white men aged 75 years or older from the third National Health and Nutrition Examination Survey (NHANES III). ⁷⁸

Excess weight, particularly central obesity, is recognized to be a major determinant of diabetes risk in all populations. Regardless of which measure of excess weight is used, the prevalence of diabetes is consistently higher among Asians than among whites at any given level of obesity. ⁷⁸ Because of this known heterogeneity of the relationship between central obesity (waist circumference) and insulin resistance, race- and ethnic-specific cut points have been developed. Race specific cut points for waist circumference in relation to insulin resistance risk are shown in Table 23-2.

It is clear that both genetic and environmental risk factors play critical roles in the development of type 2 diabetes; however, despite strong evidence of heritability, there has been little success in identifying specific genes. Numerous candidate genes have been investigated, and although some gene variants have been found that confer increased risk of type 2 diabetes, ^{79,80} many others have not been verified. No genes that explain genetic differences have been established. Because of the considerable cardiovascular risk conferred by diabetes, further studies exploring ethnic differences are essential to reduce disparities.

Inflammation

It is now recognized that atherosclerosis is an inflammatory disease. ⁸¹ Chronic, subclinical inflammation appears to be one pathophysiologic mechanism explaining the increased risk of atherosclerotic disease regardless of the amount of

Table 23—2 Recommended Waist Circumference Thresholds for the Definition of Abdominal Obesity by Race-Ethnicity

Recommended Waist Circumference

circumrerence

	Threshold for Abdominal Ob	esity	
Population	Organization	Men	Women
United States	AHA/NHLBI (ATP III)	> 102cm	> 88cm
Europidus	IDF	> 94cm	> 80cm
Middle East, MEDITERRANEAN	IDF	> 94cm	> 80cm
Sub-Saharan African	IDF	> 94cm	> 80cm
Ethnic Central and South American	IDF	> 90cm	> 80cm
Asian	IDF	> 90cm	> 80cm

AHA/NHLBI (ATP III), American Heart Association/National Heart, Lung and Blood Institute (Adult Treatment Panel III): IDF. International Diabetes Federation.

obstruction produced by that coronary disease. In the inflammatory model of atherosclerosis, it is the degree of inflammation , not the degree of obstruction, that causes acute coronary syndromes and increased CHD mortality.

Cytokines are key mediators of the inflammatory response and have been implicated in the development of atherosclerosis . Studies have shown that African Americans are more likely than white Americans to carry allelic variants demon strated to increase production of inflammatory cytokines. 82

There is now accumulating evidence that markers of subclinical inflammation may indeed predict future CHD events. One of the most studied markers is C-reactive protein (CRP), ⁸⁴ and with the use of a highly sensitive assay (hsCRP), elevation of hsCRP has been found to be associated with several major CHD risk factors and with unadjusted and age-adjusted projections of 10-year CHD risk in both men and women. ^{83,84}

Levels of hsCRP have been shown to vary by race and ethnicity. 85 In one study, with use of the NHANES data base, CRP levels were found to be highest among non-Hispanic black men and Mexican American women. According to multiple logistic regression analysis, cigarette smoking and increased age, BMI, and systolic blood pressure in men and BMI and diabetes in women were strongly associated with a greater likelihood of CRP levels > 1.0 mg/dL (P < 0.001).

Whereas hsCRP levels have been shown to be related to cardiovascular risk factors, public health approaches to modification of hsCRP levels have been less well studied. One study, 86 however, addressed physical fitness and its relation to hsCRP by ethnicity. LaMonte and colleagues hypothesized that physical fitness might protect against high levels of hsCRP. They analyzed data from a subset of 44 African American women, 45 Native American women, and 46 white women who were part of the Cross-Cultural Activity Participation Study (CAPS) in the mid-1990s.

In CAPS, physical fitness was determined by exercising on a treadmill while both speed and incline were increased, and the women continued on the treadmill until they reached their point of exhaustion. Each woman's treadmill time was adjusted for her age, and women in each of the three ethnic groups were divided into three levels of fitness (low, moderate, and high) on the basis of their treadmill tests. The researchers assessed CRP levels by race, fitness, obesity, and waist size. They found that CRP levels were 4.3 mg/L in African American women, 2.5 mg/L in Native American women, and 2.3 mg/L in white women. They also found that women with low fitness had significantly higher CRP levels (4.3 mg/L) than did those in the moderate (2.6 mg/L) and high (2.3 mg/L) fitness categories. They also reported that CRP was significantly elevated in women with the highest BMI. Women with BMI values from 18.5 to 24.9 had hsCRP levels of 1.9 mg/L, whereas overweight and obese women had hsCRP levels of 4.2 mg/L. Finally, they found that women whose waists measured more than 35 inches had CRP concentrations of 4.2 mg/L, whereas those with waist circumference of less than 35 inches had CRP levels of 2.5 mg/L. These data provide evidence of a key mechanism through which chronic stressors may accelerate atherosclerosis and may have important implications for certain racial and ethnic populations.

Obesity

Obesity is the most important cause of preventable death and the second leading cause of premature death in the United States. About 300,000 excess deaths are linked to obesity and its complications annually. Excessive adiposity is a major cause of hypertension, dyslipidemia, and type 2 diabetes mellitus. ⁸⁷ These clinical risk factors are known to be the primary precursors of CVD, and the obesity epidemic has the potential to reduce further gains in the US life expectancy, ⁸⁸ largely through an effect on cardiovascular risk and mortality. ⁸⁹ As the prevalence of obesity and overweight in the United States increases, the CVD consequences may also be concurrently increasing.

The prevalence of obesity doubled in US adults between 1980 and 2004. 90,91 Although data from 2005 to 2006 have shown no statistically significant increase in the prevalence of obesity since 2004, more than one third of US adults are now obese. 92 According to NHANES data, the increase in prevalence of overweight and obesity has been similar across racial and ethnic groups in both men and women during the past three decades; however, there are racial differences in the prevalence of obesity as well as racial differences in the prevalence of health complications associated with obesity. 93,94

Data from Behavioral Risk Factor Surveillance System surveys conducted during 2006-2008 show that African American women had the greatest prevalence of obesity (39.2%), followed by African American men (31.6%), His panic women (29.4%), Hispanic men (27.8%), non-Hispanic white men (25.4%), and non-Hispanic white women (21.8%). According to these data, African Americans (35.7%) had 51% greater prevalence of obesity, and Hispanics (28.7%) had 21% greater prevalence, compared with non-Hispanic whites (23.7%). Other studies have also shown that the prevalence of obesity among Asian Americans has been much lower than the national average, but differences between different Asian groups are considerable, and the highest prevalence of obesity has been reported among native Hawaiians and Samoans. 95

Despite obesity's disproportionately high prevalence in African Americans and Hispanic Americans, ⁹⁶ studies suggest that its adverse impact on cardiovascular risk and mortality may be reduced in some minority populations. ^{97,98} For example, even though African Americans have a higher prevalence of hypertension, diabetes, and hypercholesterolemia ⁹⁹ than white Americans do, evidence suggests that the relation ship of BMI to most CVD risk factors is steeper in whites, suggesting a stronger influence of obesity on risk factor levels in whites than in African Americans. Previous studies have suggested that for a given BMI category, rates of CVD death among African Americans are lower compared to whites. ^{100 · 103}

This observation indicates that even with higher rates of risk factors in African Americans, risk factors are more strongly associated with increasing BMI in whites. These paradoxical observations might be explained by the higher rates of type 2 diabetes mellitus, hypertension, elevated LDL-C, and low HDL among African Americans than among



whites in the normal weight group. Moreover, although the relative associations between BMI and cardiometabolic risk factors are stronger in whites than in African Americans, there is a strong association between increasing BMI and cardiometabolic risk factors in both groups.

Ø

Tuxedo

Smoking is an independent risk factor for CVD. There are a number of ways that smoking contributes to the pathogenesis of CVD. Smoking impairs lipoprotein metabolism, increases blood thrombogenicity, ¹⁰⁴ reduces the distensibility of blood vessel walls, ¹⁰⁵ and induces a proinflammatory state. ^{106,107} Smokers have higher serum levels of cholesterol and lower plasma concentrations of HDL-C. Also, cigarette smoking is associated with higher plasma concentrations of triglycerides.

According to the 2008 National Health Interview Survey, approximately 20.5% of adults are current smokers. ¹⁰⁸ Data from national surveys indicate broad disparities in cigarette smoking by race or ethnicity. Native Americans and Alaskan natives have the highest smoking prevalence (36.4%), followed by whites (21.4%), African Americans (25), Hispanics (19.8%), and Asians (13.3%). ¹⁰⁹ There are also large differences by ethnicity in receiving advice from providers to quit smoking. African Americans and Hispanics are significantly less likely to be offered assistance with cessation.

Nationally, only 21% of Hispanics report receiving regular care from a racially concordant physician, compared with 88% of whites and 23% of blacks. ¹¹⁰ The lower frequency of smoking cessation advice among Hispanics and African Americans reflects disparities in health care providers' perceptions of the need for or effectiveness of cessation advice in this ethnic subgroup.

Moreover, interest in quitting and attempts to quit differ in ethnic groups. Only 62% of Hispanic smokers reported wanting to quit compared with 71% of whites, 68% of African Americans, and 70% of Native Americans. The percentage of smokers who quit also varies sharply by ethnic group, with the highest level of success among whites (51%) and the lowest for African Americans (37%). 111

RISK CALCULATION AMONG ETHNIC GROUPS

Prediction of cardiovascular risk is a cornerstone of CVD prevention and treatment. The most commonly used risk prediction algorithm comes from the Framingham Heart Study, which has developed multivariable mathematical functions that assign weights to major CHD risk factors such as sex, age, blood pressure, total cholesterol, LDL-C, HDL-C, smoking status, and diabetes status. ¹¹²⁻¹¹⁵ The Framingham risk prediction algorithm is used to predict the risk a person free of CVD has for developing CHD within a certain period. One of the concerns about this algorithm, however, is that the Framingham Heart Study consists primarily of white middle-class individuals; thus, there is concern as to whether this algorithm can be generalized to populations of different racial and ethnic groups.

To test the validity of the Framingham CHD prediction functions in various populations, the National Heart, Lung, and Blood Institute organized a workshop for this purpose. Sexspecific CHD functions were derived from Framingham data for prediction of coronary death and myocardial infarction. These functions were applied to six prospectively studied, ethnically diverse cohorts (n = 23,424) including whites, African Americans, Native Americans, Japanese American men, and Hispanic men: the ARIC study (1987-1988), Physicians' Health Study (1982), Honolulu Heart Program (1980-1982), Puerto Rico Heart Health Program (1965-1968), Strong Heart Study (1989-1991), and Cardiovas cular Health Study (1989-1990).

The performance or ability to accurately predict CHD risk of the Framingham functions was compared with the performance of risk functions developed specifically from the individual cohorts' data. Comparisons included evaluation of the equality of relative risks for standard CHD risk factors, discrimination, and calibration. This workshop found that for white men and women and for black men and women, the Framingham functions performed reasonably well for predicting CHD events within 5 years of follow-up. Among Japanese American and Hispanic men and Native American women, the Framingham functions systematically overestimated the risk of 5-year CHD events; but after recalibration, taking into account different prevalences of risk factors and underlying rates for development of CHD, the Framingham functions worked well in these populations. ¹¹⁶

To reduce some of the limitations of risk assessment in different populations, investigators have considered adding data to the Framingham algorithm. In one study, investigators used data from the ARIC study and then validated their model in a second cohort, the NHANES linked to the National Death Index. 117 They assessed the effect of measures of socioeconomic status (SES), specifically having < 12 years of education or low income, on model discrimination and calibration when added to the Framingham risk score in a prospective cohort. They found that based on Framingham risk score alone, persons of higher and lower SES had a predicted CHD risk of 3.7% and 3.9%, respectively, compared with the observed risks of 3.2% and 5.6%. When they added SES to the model, predicted risk estimates improved to 3.1% and 5.2% for those with higher and lower SES, closely mirroring the actual observed outcomes. Inclusion of SES in the model resulted in upgrading of risk classification for 15.1%of low-SES participants (95% CI, 13.9-29.4%). 117

Other investigators have advocated the addition of measures of subclinical atherosclerosis to the Framingham algorithm to minimize ethnic discrepancies. One subclinical atherosclerosis measure of interest is measurement of coronary calcium. Several studies evaluating the prognostic accuracy of the measurement of coronary calcium by computed tomography have shown that coronary calcification is predictive of coronary events independently of standard risk factors or risk factor scores. ¹¹⁸⁻¹²¹

Data from several studies, including MESA, have determined that coronary calcium prevalence differs by race and ethnicity. In MESA, coronary calcification was measured in 6814 white, black, Hispanic, and Chinese men and women aged 45 to 84 years with no clinical CVD. ¹²² After multiple adjustments, compared with whites, the relative risks for having coronary calcification were 0.78 (95% CI, 0.74 to 0.82) in blacks, 0.85 (95% CI, 0.79 to 0.91) in Hispanics, and 0.92 (95% CI, 0.85 to 0.99) in Chinese.

Further data from MESA suggest that despite different prevalence, the predictive value of coronary calcium in various ethnic groups remains. 123 Participants in MESA were observed for a median of 3.8 years, during which time there were 162 coronary events, of which 89 were major events (myocardial infarction or death from CHD). In comparison with participants with no coronary calcium, the adjusted risk of a coronary event was 7.7-fold greater among participants with coronary calcium scores between 101 and 300 and 9.7-fold greater among participants with scores above 300 (P < 0.001 for both comparisons). Among the four racial and ethnic groups, a doubling of the calcium score increased the risk of a major coronary event by 15% to 35% and the risk of any coronary event by 18% to 39%.

Racial and ethnic minority populations in the United States bear a disproportionate burden of death and disability related to CVD. 124 The degree of racial and ethnic variations in burden of CVD goes beyond that which can be explained by risk factor variation alone. This discrepancy is thought to be partially attributable to differentials in cardiovascular care.

Disparities in cardiovascular health across the racial and ethnic groups are well documented. Limited access to care, delays in seeking cardiac care, 125 limited health literacy and education, 126-128 mistrust of the health care system by patients of color, 129,130 inadequate cultural competence of providers, 131,132 provider stereotyping of communities of color, 133 and racism of providers 134 are likely contributors to disappearances in cardiovascular care.

Financial barriers are much more likely in African American and Hispanic cardiac patients. Data from the Kaiser Family Foundation reports show that one third of Hispanics and 21% of African Americans are uninsured compared with 13% of whites, and these groups are less likely to receive preventive care. 135,136 African Americans and Hispanics are also 10% to 40% less likely to receive outpatient secondary prevention therapies for CVD. 137 Furthermore, African Americans and Hispanics appear to receive cardiovascular care at health care organizations that perform a lower volume of procedures and have higher riskadjusted mortality after coronary artery bypass graft surgery 138 and acute myocardial infarction. 139 Mortality and morbidity gaps between ethnic groups widen further in studies that look at long-term cardio vascular outcomes after hospitalizations or procedures. 140,141

ETHNIC-SPECIFIC TREATMENT ISSUES

Are Certain Groups More or Less Sensitive to **Certain Therapies?**

Physicians have known for some time that patients of different races respond differently to a variety of cardiovascular medications. A classic example is the lesser blood pressure lowering effect of angiotensin-converting enzyme inhibitors in African Americans compared with white patients, 142 but several other examples exist as well. 143,144 In discussing issues of race and ethnicity, however, it is crucial to remember that humans are essentially the same, with as great or greater variation within self-defined racial groups as across groups. 5 In fact, race and ethnicity are poor proxies for genetic variation but can potentially lend insight into differences in response to therapy, side effects related to therapy, or novel therapeutics altogether.

The issue of race-specific treatments was brought to the forefront in 2005 with the Food and Drug Administration (FDA) approval of the first cardiovascular therapeutic that was approved for a specific racial group. 145 Hydralazine-isosorbide dinitrate is a combination pill containing two commonly available vasodilators, hydralazine and isosorbide dinitrate, that had been studied more than 20 years earlier as a potential treatment of heart failure. 146,147 Whereas earlier trials had shown this combination to be less effective than angiotensin-converting enzyme inhibition for the treatment of patients with heart failure, a beneficial effect of this combination was noted in African American patients. Thus, a further study, the African-American Heart Failure Trial (A-HeFT), was initiated. 148

This trial enrolled only African American patients who were randomized to a fixed-dose combination of isosorbide and hydralazine (target daily dose, 120 mg and 225 mg, respectively) or placebo added to their existing medications. Follow-up was planned for 18 months, but the study was stopped early (mean duration of follow-up, 10 months) because of excess mortality in the placebo group (54 deaths in the placebo group versus 32 deaths in the active therapy group, for a 43% relative risk reduction). On the basis of these impressive findings, an FDA advisory panel recommended on June 16, 2005, that BiDiL be approved specifically for the treatment of heart failure in selfdefined African American patients, the group enrolled in the A-HeFT trial.

The race-specific FDA approval of Bidil was not based on a known genetic or physiological difference between the races; rather, it reflected the population in which the study was performed. In fact, there is good reason to believe that BiDil would also be effective in non-African American patients. The case of BiDil is unique in medicine, and in the future, it is likely 23 that race and ethnicity will be used less often than true indices of genetic variation.

CONCLUSION

Excess CVD and stroke morbidity and mortality are seen among various ethnic groups. This excess burden of disease is partially explained by variations in extent or implications of risk factors, but other elements such as genetic, socioeconomic, and cultural factors are also important yet poorly understood. Better understanding of the multitude of factors contributing to disparities in cardiovascular outcomes is essential to the reduction of these disparities in addition to the improvement of cardiovascular health for all.

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CHAPTER 24

Prevention of Ischemic Heart Disease in Women

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Cardiovascular disease is the

leading cause of death among women in the United States.

KEY POINTS

- Women receive fewer preventive recommendations, such as lipidlowering therapy, aspirin, and lifestyle advice, than do men with similar Framingham risk scores.
- The Framingham risk score and the Reynolds score can be used to estimate CVD risk in women.
- Dyslipidemia treatment recommendations are similar between women and men. Clinical trials demonstrate the efficacy and safety of statin dyslipidemia therapy, including more intensive treatments, for protection against major cardiovascular events in women.
- Blood pressure and diabetes treatment recommendations do not differ between women and men. Impaired glucose tolerance during pregnancy is a risk factor for future CVD in women.
- The use of aspirin for women aged 55 to 79 years is recommended when the potential benefit of a reduction in ischemic strokes outweighs the potential harm of an increase in gastrointestinal hemorrhage.
- Women should accumulate a minimum of 30 minutes of moderate-intensity physical activity (e.g., brisk walking) daily.
- Women should consume a diet rich in fruits and vegetables. whole-grain high-fiber foods, and fish at least twice a week, with limited intake of saturated fat (<7% to 10%) and cholesterol (<300 mg/ day); alcohol intake should be

limited to no more than one drink per day; sodium intake should be limited to < 2.3 g/day; and consumption of trans-fatty acids should be as low as possible.

· Hormone therapy used for contraception, management of menopause symptoms, or other clinically indicated conditions should not be used for CVD prevention.

Cardiovascular disease (CVD), the leading cause of death among women regardless of race or ethnicity, accounts for more than 500,000 deaths in the United States each year. 1 This amounts to more deaths from CVD than from lung cancer, chronic obstructive lung disease, and breast cancer combined. 2In many countries including the United States, more women than men die of CVD every year, a fact largely unknown by physicians and the lay public. ^{2,3} Typically, women cite cancer, specifically breast cancer, as a major threat to their health. Nevertheless, the annual mortality of women from CVD is twice that for all cancers combined, and almost one in two women will die of CVD compared with one in 30 of breast cancer. 4,5 Paradoxical sex differences are observed in which women have less anatomically obstructive coronary artery disease (CAD) and relatively preserved left ventricular function yet higher rates of myocardial ischemia and mortality compared with similarly aged men. 6-9 Accordingly, the term ischemic heart disease (IHD) is more appropriate for a discussion specific to women, rather than CAD or coronary heart disease (CHD). Data from the National Heart, Lung, and Blood Institute (NHLBI)-sponsored Women's Ischemia Syndrome Evaluation (WISE) and related studies involving adverse coronary

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reactivity, 10 microvascular dysfunction, 11 and plaque erosion or distal microembolization 12,13 as contributory to female-specific IHD pathophysiology. Thus, knowledge beyond an anatomical description of obstructive CAD may provide important clues to IHD risk detection and treatment for women.

AWARENESS

In a survey conducted by the American Heart Association (AHA) in 2003, only 13% of US women (7% in 1997) perceived heart disease as a major health risk. 14 Black and Hispanic women are less likely than white women to be aware that heart disease is the primary cause of death in women. Only about one third of women recall discussing heart disease risk with their physicians. 15 Women receive fewer preventive recommendations, such as lipidlowering therapy, aspirin, and lifestyle advice, than do men with similar Framingham risk scores. 3.16

Despite the availability of numerous preventive and therapeutic options, women often do not take steps to modify their cardiac risk factors; low awareness is likely to contribute to this. Although mortality from IHD has declined gradually among men since 1979 (by 30% to 50%), mortality from IHD in women has increased during the same period. ² A greater proportion of women (52%) than men (42%) with myocardial infarction (MI) die of sudden cardiac death before reaching the hospital; two thirds of women who suffer MI never completely recover. ² Since the late 1970s, hospital discharges from heart failure among women have increased at a markedly faster rate than among men. ² Thus, an under standing of even subtle differences between men and women in development and progression of IHD and in the use of proven

therapies and response to therapy could have a significant impact on improving outcomes. Even modest preventive measures can have an enormous impact. It is projected that a reduction in the death rate due to chronic diseases by just 2% during one decade would prevent 36 million deaths. ¹⁷

It has long been recognized that the first presentation with IHD occurs on average 10 years later among women than among men, usually after menopause. Although IHD, in general, is manifested earlier in less developed countries, the approximate 8- to 10-year age gap in time at onset between men and women remains universal (Fig. 24-1). The INTER HEART study, a large cohort study of more than 52,000 individuals with MI, has demonstrated that the age gap between the sexes is consistent across various socioeconomic, climatic, and cultural environments. ¹⁸ However, lifetime IHD risk for women is essentially equivalent to that of men. ¹⁹ Menopause is associated with a threefold increase in risk, although it is unclear if this is simply attributable to age. ²⁰ It

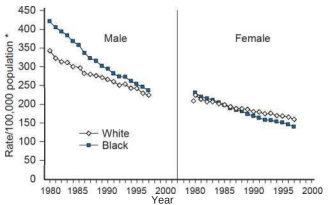


FIGURE 24-1 Death rates for coronary heart disease by sex and race in the United States from 1980 through 1997. *Rates are age adjusted to the 2000 standard. (From Cooper R, Cutler J, Desvigne-Nickens P, et al: Trends and disparities in coronary heart disease, stroke, and other cardiovascular diseases in the United States: findings of the national conference on cardiovascular disease prevention. Circulation 102:3137, 2000.)

has been widely speculated that the observed sex-related age **399** difference reflects premenopausal protection afforded by circulating estrogen; however, younger premenopausal women face a twofold increase in MI-related mortality compared with age-matched men (Fig. 24-2). ²¹ The role that endogenous and exogenous reproductive hormones play in IHD remains poorly understood.

Women are underrepresented in IHD prevention studies. Equal inclusion of women in large cardiovascular trials is now mandated by the NHLBI. ²² In practice, women are less likely to receive preventive therapy, possibly because of a 24 lack of perceived sex-specific benefit resulting from studies predominantly conducted in men. Notably, evidence-based guidelines for CVD prevention in women were published by the AHA in 2004 and later updated in 2007. ^{5,14,23}

RISK STRATIFICATION

The 2007 AHA update recommends a scheme for a general risk stratification approach to the female patient that classifies her as at high risk, at risk, or at optimal risk (Table 24-1). The 2007 AHA update focuses on the high average lifetime risk for CVD, which approaches one in two women. Conversely , the Framingham risk score focuses on a relatively narrow period of 10-year risk of CHD death or MI that does not adequately reflect the long-term or lifetime risk of women. The limitations of risk stratification with the Framingham risk score in various populations of women are well recognized . These include the lack of inclusion of family history, obesity, metabolic syndrome, and physical inactivity; inac clean estimation of risk in nonwhite populations; and the preponderance of low risk scores in women despite a high prevalence of subclinical disease. ²⁴

A Framingham risk score > 20% can identify a woman at high risk, but a lower Framingham risk score is insufficient to ensure that an individual woman is at low lifetime risk. The Reynolds risk score, derived from almost 25,000 particles in the Women's Health Study, was recently suggested as a superior alternative risk score for women, ²⁵ although validation and translation into clinical practice are needed. ²⁶ The Reynolds score incorporates novel risk markers, including high-sensitivity C-reactive protein (hsCRP) and a family

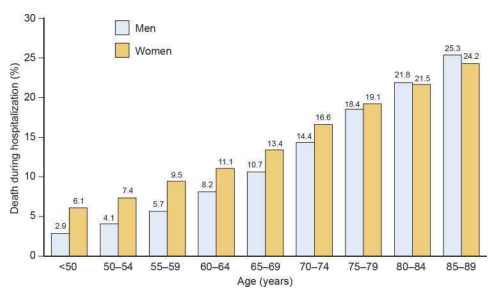


FIGURE 24-2 Sex-based differences in early mortality after myocardial infarction. (From Vaccarino V Parsons L, Every NR, et al: Sex-based differences in early mortality after myocardial infarction. National Registry of Myocardial Infarction 2 Participants. N Engl J Med 341:217, 1999.)

TABLE 24—1 Classification of Cardiovascular Disease Risk in Women

Risk Status

Criteria

High risk

Established coronary heart disease Cerebrovascular disease Peripheral arterial disease

Abdominal aortic aneurysm

End-stage or chronic renal disease Diabetes mellitus 10-year Framingham global risk > 20%††††††

At risk > 1 major risk factors for CVD, including

Cigarette smoking Poor diet

Physical inactivity
Obesity, especially central adiposity

Family history of premature CVD (CVD at < 55 years of age in male relative and < 65 years of age in female relative)

Hypertension Dyslipidemia

Evidence of subclinical vascular disease (eg, coronary

calcification) Metabolic syndrome

Poor exercise capacity on treadmill test and/or abnormal

heart rate recovery after stopping exercise

Optimal risk

Framingham global risk < 10% and a healthy lifestyle, with no risk factors

enrolled in the HERS study (1993), all of whom had established CHD, 91% failed to meet the National Cholesterol Education Program (NCEP) Adult Treatment Panel (ATP II, 1993) LDL-C goal of < 100 mg/dL. ³² In a trial in patients with stable IHD, 31% of men and only 12% of women reached LDL-C of < 100 mg/dL. ³³ In the WISE study, which enrolled women with chest pain and myocardial ischemia documented on noninvasive testing between 1996 and 2003, only 24% met their LDL-C goals. 34 These studies document less aggressive treatment of dyslipidemia in high-risk women, including those with documented IHD, compared with men. 4,5,29

RISK FACTORS AND RISK REDUCTION INTERVENTIONS

Age

Age is one of the most powerful risk factors for the development of IHD and accompanying clinical events. IHD events including MI lag 10 to 15 years in women compared to men. As men age, they continue to have a linear increase in IHD incidence, particularly during the fifth and sixth decades, whereas women have an almost exponential increase in IHD incidence after the age of 60 years, ³⁵ although population-adjusted curves suggest that this may be due to survivor effects. One in eight women between the ages of 45 and 64 years has evidence of CAD, which increases to one in three after the age of 65 years. ³⁶ With more women living to elderly ages, a majority of IHD victims are not surprisingly now women. It is for these reasons that NCEP ATP III has different age cutoffs for sex; at or above 55 years is the age considered a risk factor for women compared with 45 years for

Family History

It is well established that CVD has a hereditary component. However, few prospective studies have evaluated the relationship between family history and future IHD events in

††††††Or at high risk on the basis of another population-adapted tool used to assess global risk.

From Mosca L, Banka CL, Benjamin EJ, et al: Evidence-based guidelines for cardiovascular disease prevention in women: 2007 update. Circulation 115:1481, 2007.

history of CAD, that may aid in unique detection of risk in women. It also has the advantage of including cerebrovascular events as an outcome; this point is important because of the higher frequency of stroke in women compared to men. Application of the Reynolds risk score reclassified 40% to 50% of women with an intermediate Framingham risk score into higher or lower risk categories. ²⁷

D'Agostino and colleagues ²⁸ proposed a sex-specific multivariable risk factor algorithm that can be used to assess general CVD risk and risk of individual cardiovascular events (coronary, cerebrovascular, and peripheral arterial disease and heart failure). The authors used Cox proportional hazards regression to evaluate the risk for development of a first car diovascular event in 8491 Framingham study participants (mean age, 49 years; 4522 women) who attended a routine examination between 30 and 74 years of age and were free of CVD. Sex-specific multivariable risk functions ("general CVD" algorithms) were derived that incorporated age, total and high-density lipoprotein cholesterol, systolic blood pressure, treatment of hypertension, smoking, and diabetes status. During 12 years of follow-up, 1174 participants (456 women) developed a first cardiovascular event. The general CVD algorithm demonstrated good discrimination (C-statistic, 0.763 [men] and 0.793 [women]) and calibration.

There is considerable evidence that elevated levels of total cholesterol (TC) and low-density lipoprotein cholesterol (LDL-C) increase IHD risk in women, ^{4.29} although these relationships are less prominent in elderly women compared with younger women. ^{4.30} In contrast, low levels of high- density lipoprotein cholesterol (HDL-C) and elevated levels of triglyceride (TG) impart comparable risk in young and elderly women. ^{4.30} Despite this evidence, men are more likely to have their lipid levels measured, and abnormal values are generally treated more aggressively than in women. ^{4.31} In women

women. Retrospective review of much larger data bases of women (eg, the Nurses' Health Study) has demonstrated conflicting results. In women with a family history of MI before the age of 60 years, the age-adjusted relative risk was 2.8 for nonfatal MI and 5.0 for fatal cardiovascular events. ³⁷ With adjustment for other cardiovascular risk factors, family history remained an independent risk factor for CVD. NCEP ATP III defines family history of premature CHD before the age of 65 years for women and before the age of 55 years for men. ³⁸ Although family history of premature CHD signifies high risk, risk remains elevated when the definition is broadened to include older first-degree relatives.

Dyslipidemia

More than half of the women in the United States have a TC level > 200 mg/dL, and 36% have LDL-C concentration > 130 mg/dL. ² Notably, only 13% of women have HDL-C concentration < 40 mg/dL (compared with 23% of Americans overall); the higher threshold of < 50 mg/dL is used for women. The relative risk for IHD events associated with elevation of various lipid variables was determined in a nested case-control study from the Nurses' Health Study. Among 32,826 healthy women, the multivariable relative risks (adjusted for hsCRP, homocysteine, and other traditional cardiac risk factors) for the highest quintiles of lipid variables were as follows: apolipoprotein B: RR, 4.1 (2-8.3); low levels of HDL-C: RR, 2.6 (1.4-5); LDL-C: RR, 3.1 (1.7-5.8); and TG: RR, 1.9 (1-3.9). ³⁹ Adverse changes in lipid profile accompany menopause. ⁴⁰ Perimenopausal TG levels are the

TABLE 24-2 Major Lipid-Lowering Trials Using Statin Therapy for Coronary Heart Disease (CHD) Prevention						
Study	Patients (n)	Women, n (%)	Prevention Category	Drug	Year	Risk Reduction in Major CHD Events in Women (%)
4S ⁴³	4444	827 (19)	Secondary	Simvastatin	1994	35
CARE44	4159	576 (14)	Secondary	Pravastatin	1996	46
LIPID45	9014	1516 (17)	Secondary	Pravastatin	1998	11
AFCAPS/TexCAPS' 46	6605	997 (15)	Primary	Lovastatin	1998	46
PROSPER47	5804	3000 (52)	Both	Pravastatin	2002	NS benefit for women
HPS ⁴⁸	20,536	5082 (25)	Primary	Simvastatin	2002	19
ALLHAT-LLT ⁴⁹	10,355	5051 (49)	Primary (hypertension) 14% CHD	Pravastatin	2002	Sex-specific data not reported; NS in total cohort
GREACE ⁵⁰	1600	344(21)	Secondary	Atorvastatin	2002	54
ASCOT-LLA ₅₁	10,305	1942 (19)	Primary (hypertension)	Atorvastatin	2003	NS benefit for women

NS. Not significant.

From Lavie CJ, Wenger NK: Special patient populations: women and elderly. In Ballantyne CM, editor: Clinical lipidology, Philadelphia, 2009, Saunders/Elsevier, p 465.

most erratic but follow roughly the same pattern of increased TC and LDL-C on average by $\sim \! 10\%$ from levels 6 months before menopause. Menopause can influence HDL-C less dramatically, with mild declines noted.

Dyslipidemia is strongly predictive of IHD risk in older women. Women in the Framingham study with TC > 265 mg/dL were at two to three times greater risk for experiencing an IHD event compared with women with TC < 205 mg/dL. Overall, a 1% increase in TC translated to a 2% increase in IHD incidence. 41 Elevated TG levels and low HDL-C levels are more closely associated with the risk of IHD in women, especially in those at least 65 years old. Lower levels of HDL-C are associated with an increased risk of IHD in the Lipid Research Clinics Follow-up Study. HDL-C was second to age in predicting IHD risk. The ratio of TC and HDL-C was a more accurate marker of IHD risk than either level alone. The NCEP guidelines call for measurement of TC as well as of HDL-C as part of the initial cholesterol screening after the age of 20 years. The TC/HDL-C ratio is typically lower in women than in men through middle age but then increases and parallels that of men by 75 years of age. After the age of 55 years, elevated TC and TC/HDL-C ratio significantly increase IHD risk.

The protective role of lipid-lowering therapy in women, especially with the 3-hydroxy-3-methylglutaryl coenzyme A reductase inhibitors (ie, statins), is well established by numerous randomized clinical trials noting consistent reductions in CVD risk ranging from 11% to 54% 4,42-51 (Table 24-2). Although the representation of women in most trials is relatively low (averaging < 20% of the population), a positive trend for benefit was consistent across all trials. Importantly, the protective effect of statins based on these trials appeared equivalent to or greater than that observed for men. The reduction in risk of major cardiovascular events in women ranged from 11% in the Longterm Intervention with Pravastatin Ischemic Disease (LIPID) study 45 to 54% in the Greek Atorvastatin and Coronary-heartdisease Evaluation (GREACE) trial. 50 There was no significant effect in the subgroup of women in the Prospective Study of Pravastatin in the Elderly at Risk (PROSPER) 47 or in the lipidlowering arm of the Anglo-Scandinavian Cardiac Outcomes Trial (ASCOT). 51 There was no significant effect in the total cohort (men or women) in the lipid-lowering arm of the Antihypertensive and Lipid-Lowering Treatment to Prevent Heart Attack Trial (ALLHAT) 49; however, Treating to New Targets (TNT), a trial of high-dose (80 mg) versus low-dose (10 mg) atorvastatin, showed equal protection against major cardiovascular events with more intensive therapy in women,

statistically similar to that of men, without a sex difference in serious adverse events. There was modestly higher frequency of discontinuation of statins (10% versus 6.5%) and liver function test abnormalities (2.5% versus 1%) in women than in men. 52 Likewise, in patients with acute coronary syndromes (n = 4162 patients, 22% women), more intensive lipid treatment with atorvastatin 80 mg versus pravastatin 40 mg also yielded similar benefits for women and men. 53

In aggregate, these trials demonstrate the efficacy and safety of statin therapy, including more intensive treatments, for protection against major cardiovascular events in women. In addition to significant reductions of LDL-C and moderate improvements in HDL-C and TG, other potential beneficial effects of statins include improvements in endothelial function, plaque stabilization, anti-inflammatory effects, and antiplatelet effects, among others. Higher risk patients, such as secondary prevention, offer the best short-term efficacy targets. Whereas it has been argued that treatment of dyslipidemic young women is not cost-effective, recent prevention guidelines focused on longer term lifetime risk question this rationale. 5,25,27 In addition, women carry a high short-term mortality rate associated with first IHD events (more than twice that for men), providing further support for consideration of statin therapy for both primary and secondary prevention of IHD in women (Table 24-

On the basis of a number of trials in high-risk subjects, ^{53,54} the 2004 revision of the NCEP ATP III guidelines introduced a new LDL-C goal of < 70 mg/dL for patients at very high risk, which includes women with established CVD and multiple other cardiovascular risk factors. 55 A meta-analysis of primary prevention trials (excluding the recent JUPITER trial) 56 concluded that pharmacological lipid lowering, primarily with statins, does not reduce mortality or events in women without known CVD. 57 Caution, however, is warranted in the interpretation of this finding because of the relatively small number of cardiovascular events and the relatively young age of the female enrollees. As young women have a lower absolute risk than that of men in a limited observational period, it is conceivable that the number needed to treat is significantly higher for women than for men. As statins primarily affect LDL-C reduction, one can speculate whether targeting of HDL-C or TGs may be more promising in women compared to men. The documented benefits of lipid-lowering therapy in high-risk women, on the other hand, are convincing and robust. It is therefore important to implement aggressive pharmacotherapy to new lipid targets in high-risk women. 58



TABLE 24—3 Guidelines for Prevention of Cardiovascular Disease in Women: Clinical Recommendations

Lifestyle Interventions

Cigarette Smoking

Women should not smoke and should avoid environmental tobacco smoke. Provide counseling, nicotine replacement, and other pharmacotherapy as indicated in conjunction with a behavioral program or formal smoking cessation program (Class I, Level B).

Physical Activity

Women should accumulate a minimum of 30 minutes of moderate-intensity physical activity (eg, brisk walking) on most and preferably all days of the week (Class I, Level B)

Women who need to lose weight or sustain weight loss should accumulate a minimum of 60 to 90 minutes of moderate-intensity physical activity (eg, brisk walking) on most and preferably all days of the week (Class I, Level C).

Rehabilitation

A comprehensive risk reduction regimen, such as cardiovascular or stroke rehabilitation or a physician-guided home- or community-based exercise training program, should be recommended to women with a recent acute coronary syndrome or coronary intervention, new-onset or chronic angina, recent cerebrovascular event, peripheral arterial disease (Class I, Level A), or current/prior symptoms of heart failure and LVEF < 40% (Class I, Level B).

Dietary Intake

Women should consume a diet rich in fruits and vegetables; choose whole-grain, high-fiber foods; consume fish, especially oily fish,* at least twice a week; limit intake of saturated fat to < 10% of energy and if possible to < 7%, cholesterol to < 300 mg/day, alcohol intake to no more than 1 drink per day, and sodium intake to < 2.3 g/day (approximately 1 tsp salt). Consumption of *trans-fatty* acids should be as low as possible (eg, < 1% of energy) (Class I, Level B).

Weight Maintenance or Reduction

Women should maintain or lose weight through an appropriate balance of physical activity, calorie intake, and formal behavioral programs when indicated to maintain or to achieve a BMI between 18.5 and 24.9 kg/m and a waist circumference < 35 inches (Class I, Level B).

Omega-3 Fatty Acids

As an adjunct to diet, omega-3 fatty acids in capsule form (approximately 850 to 1000 mg of EPA and DHA) may be considered in women with CHD, and higher doses (2 to 4 g) may be used for treatment of women with high triglyceride levels (Class Ilb, Level B).

Depression

Consider screening women with CHD for depression and refer or treat when indicated (Class Ila, Level B)

Major Risk Factor Interventions

Blood Pressure—Optimal Level and Lifestyle

Encourage an optimal blood pressure of < 120/80 mm Hg through lifestyle approaches such as weight control, increased physical activity, alcohol moderation, sodium restriction, and increased consumption of fresh fruits and vegetables and low-fat dairy products (Class I, Level B).

Blood Pressure—Pharmacotherapy

Pharmacotherapy is indicated when blood pressure is > 140/90 mm Hg or at an even lower blood pressure in the setting of chronic kidney disease or diabetes (> 130/80 mm Hg).

Thiazide diuretics should be part of the drug regimen for most patients unless contraindicated or if there are compelling indications for other agents in specific vascular diseases.

Initial treatment of high-risk women should be with beta blockers and/or ACE inhibitors or ARBs, with the addition of other drugs such as thiazides as needed to achieve goal blood pressure (Class I. Level A).

Lipid and Lipoprotein Levels—Optimal Levels and Lifestyle

The following levels of lipids and lipoproteins in women should be encouraged through lifestyle approaches: LDL-C < 100 mg/dL, HDL-C > 50 mg/dL, triglycerides < 150 mg/dL, and non-HDL-C (total cholesterol minus HDL cholesterol) < 130 mg/dL (Class I, Level B). If a woman is at high risk or has hypercholesterolemia, intake of saturated fat should be < 7% and cholesterol intake < 200 mg/day (Class I, Level B).

Lipids—Pharmacotherapy for LDL Lowering, High-Risk Women

Use LDL-C-lowering drug therapy simultaneously with lifestyle therapy in women with CHD to achieve an LDL-C < 100 mg/dL (Class I, Level A) and similarly in women with other atherosclerotic CVD or diabetes mellitus or 10-year absolute risk > 20% (Class I, Level B).

A reduction to < 70 mg/dL is reasonable in very-high-risk women swith CHD and may require an LDL-lowering drug combination (Class IIa, Level B).

Lipids—Pharmacotherapy for LDL Lowering, Other At-Risk Women

Use LDL-C-lowering therapy if LDL-C level is > 130 mg/dL with lifestyle therapy and there are multiple risk factors and 10-year absolute risk 10% to 20% (Class I, Level B).

Use LDL-C-lowering therapy if LDL-C level is > 160 mg/dL with lifestyle therapy and multiple risk factors even if 10-year absolute risk is < 10% (Class I, Level B).

 $\begin{tabular}{ll} Use LDL-C-lowering the rapy if LDL > 190 mg/dL regardless of the presence or absence of other risk factors or CVD on lifestyle the rapy (Class I, Level B) . \\ \end{tabular}$

Lipids—Pharmacotherapy for Low HDL or Elevated Non-HDL, High-Risk Women

Use niacin or fibrate therapy when HDL-C is low or non-HDL-C is elevated in high-risk women after LDL-C goal is reached (Class IIa, Level B).

Lipids—Pharmacotherapy for Low HDL or Elevated Non-HDL, Other At-Risk Women

I consider niacin or fibrate therapy when HDL-C is low or non-HDL-C is elevated after LDL-C goal is reached in women with multiple risk factors and a 10-year absolute risk 10% to 20% (Class Ilb, Level B).

Diabetes Mellitus

Lifestyle and pharmacotherapy should be used as indicated in women with diabetes (Class I, Level B) to achieve HbA1c < 7% if this can be accomplished without significant hypoglycemia (Class I, Level C).

Preventive Drug Interventions

Aspirin, High Risk

Aspirin therapy (75 to 325 mg/day) - should be used in high-risk *** women unless contraindicated (Class I, Level A). If a high-risk ** woman is intolerant of aspirin therapy, clopidogrel should be substituted (Class I, Level B).

Aspirin-Other At-Risk or Healthy Women

In women > 65 years of age, consider aspirin therapy (81 mg daily or 100 mg every other day) if blood pressure is controlled and benefit for ischemic stroke and MI prevention is likely to outweigh risk of gastrointestinal bleeding and hemorrhagic stroke (Class Ila, Level B) and in women < 65 years of age when benefit for ischemic stroke prevention is likely to outweigh adverse effects of therapy (Class Ilb, Level B).

Beta Blockers

Beta blockers should be used indefinitely in all women after MI, acute coronary syndrome, or left ventricular dysfunction with or without heart failure symptoms, unless contraindicated (Class I, Level A).

ACE Inhibitors/ARBs

ACE inhibitors should be used (unless contraindicated) in women after MI and in those with clinical evidence of heart failure or LVEF < 40% or with diabetes mellitus (Class I, Level A). In women after MI and in those with clinical evidence of heart failure or LVEF < 40% or with diabetes mellitus who are intolerant of ACE inhibitors, ARBs should be used instead (Class I, Level B).

Aldosterone Blockade

Use aldosterone blockade after MI in women who do not have significant renal dysfunction or hyperkalemia who are already receiving therapeutic doses of an ACE inhibitor and beta blocker and have LVEF < 40% with symptomatic heart failure (Class I, Level B).

A 2003 national survey highlighted that women remain undertreated compared with men and that significantly fewer high-risk women achieved their LDL-C goal compared with men. 59

From the recent JUPITER trial, 17,802 apparently healthy men and women with LDL-C levels of $< 130 \, mg/dL$ and hsCRP levels of > 2.0 mg/L were randomized to 20 mg of rosuvastatin daily versus placebo. 56 The trial was stopped early after a median follow-up of 1.9 years, with rosuvastatin reducing LDL-C levels by 50% and hsČRP levels by 37%. Rosuvastatin significantly reduced the primary endpoint, a composite of nonfatal MI, stroke, hospitalization for unstable revascularization, and CVD death, by 44% compared with individuals treated with placebo. This reduction was observed across nearly all of the endpoints, including a 55% reduction in nonfatal MI, a 48% reduction in the risk of nonfatal stroke, and a 47% reduction in the risk of "hard" cardiac events (a composite of MI, stroke, and CVD death). Consistent effects were observed in all subgroups including women. Among the 6801 women (38%) included in JUPITER, rosuv-astatin significantly reduced the primary composite end point by 46%, a magnitude similar to that observed in men.

Women are often concerned about stroke more than MI. Data from statin trials such as JUPITER, which showed a 48% reduction in the risk of nonfatal stroke with rosuvastatin, might help physicians provide a more convincing argument for women who may be reluctant about taking statins, especially if familial risk of stroke is great relative to that of MI.

Tuxedo

Cigarette smoking is the most preventable IHD risk factor in women and men and increases the risk of MI in women relatively

greater than in men. ^{60,61} More than 60% of MIs in women younger than 50 years are attributable to smoking, as are 21% of all deaths from IHD. For female smokers, a dose- dependent relationship between consumption and risk has been described. ^{62,63} Rosenberg and colleagues ⁶² cited an increased risk of nonfatal MI from 2.4 in women who smoke 15 to 24 cigarettes/day to 7 for those who smoke > 25 cigarettes/day. In addition, smoking increases the IHD risk when other cardiac risk factors are present or when oral contraceptive pills are being taken. Women not taking oral contraceptive pills who smoke > 25 cigarettes/day have a 4.8-fold relative risk of nonfatal MI compared with a 23-fold relative risk in female smokers taking oral contraceptives.

In the United States, fewer women than men currently smoke, and the prevalence of smoking among women has declined during the past three decades. However, tobacco use has decreased more dramatically in men than in women. More worrisome is the increased prevalence of female adolescents who are smoking. Compared with male smokers, women more often smoke to relieve stress, anger, boredom, or depression. Women are more likely than men to cite smoking as a strategy for weight loss, and they more often give weight gain as a major reason for relapsing. ⁶⁴ All women benefit from smoking cessation, regardless of age at cessation, which should provide a tremendous incentive for elderly smokers to quit. In the Nurses' Health Study, women who

ACE, angiotensin-converting enzyme; ARB, angiotensin receptor blocker; BMI, body mass index; CHD, coronary heart disease; CVD, cardiovascular disease; DHA, docosahexaenoic acid; EPA, eicosapentaenoic acid; HDL-C, high-density lipoprotein cholesterol; LDL-C, low-density lipoprotein cholesterol; LVEF, left ventricular ejection fraction; MI, myocardial inferention.

^{*}Pregnant and lactating women should avoid eating fish potentially high in methylmercury (eg, shark, swordfish, king mackerel, or tile fish) and should eat up to 12 oz/wk of a variety of fish and shellfish low in mercury and check the Environmental Protection Agency and the US Food and Drug Administration's websites for updates and local advisories about the safety of local catch.

[·] A drink equivalent is equal to a 12-oz bottle of beer, a 5-oz glass of wine, or a 1.5-oz shot of 80-proof spirit.

ssssss Criteria for high risk include established CHD, cerebrovascular disease, peripheral arterial disease, abdominal aortic aneurysm, end-stage or chronic renal disease, diabetes mellitus, and 10-year Framingham risk >20%.

[^]Criteria for very high risk include established CVD plus any of the following: multiple major risk factors, severe and poorly controlled risk factors, diabetes mellitus. || Dietary supplement niacin should not be used as a substitute for prescription niacin.

After percutaneous intervention with stent placement or coronary artery bypass grafting within the previous year and in women with noncoronary forms of CVD, use current quidelines for aspirin and clopidogrel.

From Mosca L. Banka CL. Benjamin EJ, et al: Evidence-based guidelines for cardiovascular disease prevention in women: 2007 update. Circulation 115:1481, 2007.

404 stopped smoking experienced an immediate benefit as well as a further long-term decline in IHD risk to levels comparable to those of women who never smoked. ⁶⁵ Total mortality risk among former smokers decreases nearly to that of never-smokers within 10 to 14 years of cessation. ⁶⁶ The risk of an index MI in women declines soon after the cessation of smoking and is largely dissipated after 2 to 3 years. ⁶⁷

Compared with men, women find it more difficult to quit smoking with and without treatment and are also more likely to relapse. Simple encouragement to stop smoking 24 hours a day should be part of every follow-up encounter with an active smoker. In women who are unable to quit without assistance, use of nicotine replacement and other pharmacotherapy (eg, bupropion, varenicline) in conjunction with behavioral therapy or a formal smoking cessation program is warranted.

Physical Inactivity

Sedentary lifestyle is a common risk factor for IHD in women and men. Data from the National Center for Health Statistics indicate that 39% of white women and 57% of women of color do not exercise regularly. 68 Rates of physical inactivity are highest among poor women. Increased levels of physical activity are associated with lower blood pressure, lower cholesterol levels, improved glucose metabolism, higher bone density, and improved mental health variables. Physical inactivity contributes to obesity and is an independent risk factor for MI. 69 Even modest exercise has been strongly associated with risk reduction in observational studies. Investigators in the Nurses' Health Study 70 found that 30 to 45 minutes of walking three times weekly reduces the risk of MI by 50% in women across the ages. Exercise has also been found to reduce the risk of type 2 diabetes mellitus, even in obese women and those with a family history of diabetes. 71

Physical activity has an even more beneficial role in women after MI or coronary artery bypass surgery. However, despite improved health outcomes associated with cardiac rehabilitation, fewer women than men enroll. A variety of reasons, including referral bias, difficulty with transportation, and elder and child care responsibilities, often prevent women from participating in cardiac rehabilitation programs. According to the 2007 prevention guidelines for women, women should accumulate a minimum of 30 minutes of moderate-intensity physical activity (eg, brisk walking) daily (Class I recommendation) (see Table 24-3).

Diet

In the Nurses' Health Study, the dietary score, based on intake of cereal fiber, marine-3 fatty acids, folate, glycemic load, and ratio of polyunsaturated to saturated fat, acted as an independent factor for IHD. ⁶⁹ A second report from the Nurses' Health Study evaluated the relationship between diet and cardiovascular events during 1980-1994 in 85,941 apparently healthy women. Improvement in dietary score consisting of reduced consumption of red meat, trans- fats, and high-fat products during the study was associated with a significant 16% decline in IHD incidence. 72 Current recommendations suggest that women should consume a diet rich in fruits and vegetables, whole-grain high-fiber foods, and fish at least twice a week, with limited intake of saturated fat (< 7% to 10%) and cholesterol (< 300 mg/day). Alcohol intake should be limited to no more than one drink per day and sodium intake to < 2.3 g/day. Consumption of *trans* -fatty acids should be as low as possible (see Table 24-3). Nutrition intervention trials have demonstrated

reductions in CVD and mortality, ⁷³⁻⁷⁵ although sexstratified trial results are typically not available.

Obesity

Between 1988 and 1991, the Third National Health and Nutrition Examination Survey (NHANES III) reported that one third of US adults (35% of the women and 31% of the men) were overweight. 76,77 Obesity was defined by a body mass index (BMI) of 27.8 kg/m² in men and 27.3 kg/m² in women (120% of desirable weight). Obesity is associated with glucose intolerance (or prediabetes mellitus), hypertension, and hypercholesterolemia and is also considered an independent risk factor for IHD. 76,78,79 In the Nurses' Health Study, obesity was strongly associated with an increased incidence of car diovascular events even after controlling for older age and smoking. 78 Women who were > 30% of ideal body weight had an increased risk ratio of 3.3 for nonfatal MI and IHD death. Recent data suggest that regional fat distribution, particularly a waist-to-hip ratio of > 0.88 (android body type), may be more of an accurate marker for chronic ischemic IHD risk than obesity. Several studies have reported an increased risk of MI and IHD or CVD death in women with increased truncal adiposity even after adjustment of other cardiac risk factors. 80,81

Data from the Framingham study indicate a doubling in the incidence of diabetes for both sexes during the past 30 years; most of this increase occurred in individuals with a BMI of 30 kg/m² and above. ⁸² The Coronary Artery Risk Development in Young Adults (CARDIA) study illustrated that young adults who maintained a stable BMI over time had minimal progression of risk factors and lower incidence of metabolic syndrome, regardless of baseline BMI. However, among 1358 men and 1321 women, only 16.3% maintained a stable BMI, and 73.9% had an increased BMI. 83 The increasing prevalence of obesity and diabetes in the population, despite a reduction of dietary fat intake during the past decades, highlights the importance of dietary recommendations for exercise and carbohydrate and fat intake. The Nurses' Health Study reported a positive association between dietary glycemic load and IHD in women, which was even more pronounced in overweight and obese women. 84 Thus, reducing the glycemic load in the diet should be as much of a priority as replacing saturated fats by unsaturated fats. Avoidance of refined sugars and restriction of sugar-sweetened beverages may particularly help reduce dietary glycemic load. 85 A diet high in fruits and vegetables may reduce the risk of diabetes; however, high consumption of fruit juices may be associated with an increased risk of diabetes among women. 86

Diabetes and the Metabolic Syndrome

Diabetes is a relatively greater risk factor in women than in men, increasing IHD risk by threefold to sevenfold in diabetic women compared with a twofold to threefold increase in risk for diabetic men. ²³ Furthermore, the risk of IHD death is higher in diabetic women than in diabetic men. 87 Whereas angiographic CAD is generally more prevalent in men than in women, diabetes eliminates this difference by increasing the disproportionately in women. Large-scale prospective studies, including the Framingham study and the Nurses' Health Study, report diabetes mellitus as an independent risk factor; diabetic women were six to seven times more likely to experience nonfatal MI and CVD death. In addition, the risk of CVD morbidity and mortality in women with diabetes is minimally affected by the duration of diabetes, and a significant risk for cardiovascular events is still noted in patients with diabetes of less than 4 years in duration. 41

Diabetic women have not experienced a decrease in IHD mortality during the past three decades compared with their male counterparts. ⁸⁸ In fact, there was an increase in IHD mortality in the subgroup of diabetic women. ⁸⁹ As diabetes is considered a CHD risk equivalent by the NCEP, all diabetics

women are classified as high or very high risk. A history of diabetes Hypertension in women was associated with a 37% increased IHD-related mortality. 90 Diabetes appears to have a greater adverse effect on TG, HDL-C, and LDL-C concentrations in diabetic women than in diabetic men. 91

Milder forms of glucose intolerance and asymptomatic hyperglycemia may still place women at risk for IHD events. Type 2 diabetes or milder forms of glucose intolerance tend to cluster in women with visceral obesity, hypertension, and dyslipidemia. 92 This clinical pattern confers a much higher IHD risk in diabetic women, depending on the accompanying risk factor profile. Because other risk factors, such as hypertension, smoking, and obesity, act synergistically with diabetes, control of these other risk factors attenuates the risk of MI in diabetic women.

According to the American Diabetes Association, diabetes screening should be considered for women and men older than 45 years and repeated every 3 years, if results are normal. 93 Women with a history of gestational diabetes or polycystic ovarian syndrome should be screened earlier. History of gestational diabetes doubles the risk of diabetes within 4 months postpartum, and it remains a lifelong risk factor for the development of diabetes, which is largely a risk of type 2 diabetes. 94 Fasting plasma glucose levels of > 121 mg/ dL during pregnancy increase the risk for diabetes in the early puerperium by 21-fold. 95 Expert panels recommend fasting glucose testing or oral glucose tolerance tests 6 to 12 weeks postpartum, then every 1 to 2 years in women with gestational diabetes. To prevent the development of diabetes in high-risk women, pre-pregnancy weight should be reached within 6 to 12 months postpartum, and physical activity should be recommended.

Furthermore, it is reasonable to screen for diabetes earlier if obesity, hypertension, and dyslipidemia are present because these characteristics frequently co-occur as the metabolic syndrome. The prevalence of metabolic syndrome is similar for both sexes after adjustment for age. A hallmark of the metabolic syndrome is insulin resistance, defined as impaired fasting glucose (100 to 125 mg/dL), which is considered to be a prediabetic state. For women in the Framingham study, impaired fasting glucose was associated with increased IHD risk to a similar degree as established diabetes, a finding that was not seen in the men. 96

Clinical trials demonstrate the value of risk factor management in diabetics. The Collaborative Atorvastatin Diabetes Study (CARDS) trial, which used atorvastatin 10 mg every day versus placebo in more than 3000 diabetic patients, was terminated early because of a 37% risk reduction with the statin. 97 In the Diabetes Control and Complications Trial, type 1 diabetics receiving conventional therapy observed for 6.5 years had fewer microvascular and neurological complications with no impact on major cardiovascular events 98; however, in the follow-up trial, Epidemiology of Diabetes Interventions and Complications (EDIC), 99 a 42% risk reduction for any cardiovascular event and 57% reduction for major cardiovascular events (including fatal or nonfatal MI or stroke) was observed in the intensively treated subjects. In addition, the PROspective pioglitazone Clinical Trial in macroVascular Events (PROactive) 100 study of 5238 type 2 diabetics, which randomized patients to the PPAR y agonist pioglitazone or placebo, produced a nonsignificant reduction in the primary endpoint (all-cause mortality, nonfatal MI, stroke, leg Novel Risk Factors amputation, and cardiovascular or leg revascularization). There Traditional risk factors and the Framingham risk score under was, however, a significant 20% reduction in the secondary endpoint of all-cause mortality, nonfatal MI, and stroke. Finally, the improve risk detection. 11,119-121 Women have, on average, higher Diabetes Prevention Program demonstrated efficacy with both hsCRP measures compared with men, a sex difference apparent lifestyle interventions and metformin; high-risk individuals who at the time of puberty. 122 This difference in inflammatory exercised to goal had the lowest development of diabetes, whereas markers is consistent with the higher frequency of the addition of metformin significantly reduced diabetes development by 31% in those

receiving the drug. 101 Overall, these intervention trial results 405 aimed at CVD risk in diabetes and metabolic syndrome subjects are not available stratified by sex.

The overall prevalence of hypertension is higher among women than among men, although it varies by age. Until the age of 45 years, the prevalence of hypertension is higher in men than in women. In older individuals, hypertension is more prevalent in women, with a much higher prevalence in 24 women aged 65 years and older. 102 Hypertension is two to three times more common in women taking oral contraceptives. 103 Of note, incidence rates for black individuals are approximately twice the rates for white women and men across the ages. 104

The association of hypertension and IHD has been reported in many prospective studies in both men and women. Hypertension is also the leading risk factor for congestive heart failure. The Systolic Hypertension in the Elderly Program evaluated pharmacological intervention with a thiazide diuretic in the treatment of systolic hypertension. This study included a study population of 57% women and demonstrated reductions in stroke and IHD incidence by 36% and 25%, respectively. 105 A second trial tested the benefit of beta blockers and thiazide diuretics in decreasing target organ damage from hypertension and demonstrated similar risk reduction in women and men; however, in absolute terms, the benefit in women was seen primarily for stroke, whereas in men, treatment prevented as many IHD events as strokes. 106

Blood pressure treatment recommendations, based on the Seventh Report of the Joint National Committee on the -Prevention, Detection, and Treatment of High Blood Pressure (JNC 7), do not differ between women and men. 103 An optimal blood pressure of below 120/80 mm Hg should be encouraged through lifestyle modifications, such as weight control, physical activity, alcohol moderation, sodium restriction, and increased consumption of fruits and vegetables and low-fat dairy products (see Table 24-3). Pharmacotherapy is indicated if the blood pressure remains higher than 140/90 mm Hg or 130/80 mm Hg in women with diabetes or chronic kidney disease. Once medications are initiated, thia zide diuretics should be part of the regimen unless contraindicated or if another drug has a compelling indication and achieves sufficient blood pressure control. 49 Thiazide diuretics have a favorable effect on calcium excretion and thus may be especially appropriate for use in women who are at risk for osteoporosis. Despite similar treatment rates throughout different age groups, hypertension control is especially poor in older women, with only 29% of hypertensive women aged 70 to 79 years reaching their blood pressure target. 107

Although some studies have shown that the treatment of hypertension among women conferred relatively less benefit against cardiac events compared with antihypertensive treatment among men, 106,108-111 the national guidelines recommend the same approach for treatment of hypertensive men and women. A recent publication based on data from the National Nutrition and Health Examination Survey 1999-2004 showed that hypertensive women are significantly more likely to be treated than are men but less likely to have achieved blood pressure control. 112

estimate IHD risk in women, 113-118 and novel risk markers may

arthritis and systemic lupus erythematosus, observed in women gastrointestinal bleeding. There fore, weighing of the individual compared with men 123 and suggests a more prominent role for risk of gastrointestinal or cerebral hemorrhage versus benefit in

hsCRP, are related to other IHD risk markers, such as the although the validity of this phenomenon remains unknown. metabolic syndrome, type 2 diabetes, and heart failure. 126,129-130

assessment in women. 131-133

Antiplatelet Therapy

An overview of randomized trials from the Antiplatelet Trialists' Collaboration found that aspirin (75 to 162 mg/day) was beneficial in women for secondary prevention of IHD. 134 Among patients with an acute coronary syndrome, aspirin should be part of preventive management for women and men.

Although the benefits of aspirin therapy in reducing the risk of MI, stroke, and vascular death among men and women with REPRODUCTIVE HORMONES preexisting CVD are well established, ¹³⁴⁻¹³⁶ the role of aspirin in Postmenopausal Hormone Therapy primary prevention in women is less established, primarily because of a lack of inclusion of women in clinical trials. Aspirin has proven Until the mid-1990s, our knowledge of IHD risk and postbenefit and is recommended for all women at high risk without menopausal hormone therapy was based on observational trials contraindications and for select at-risk women 5 (see Table 24-3). that consistently reported a 40% to 50% reduction in CVD. 144 In prevention of IHD in women. Earlier recommendations were based trials using estrogen alone and those using estrogen and Hypertension Optimal Treatment (HOT) study, men (53%) and question. 146 More than 2700 women were randomized to receive women (47%) were randomized to receive 75 mg of aspirin per day conjugated equine estrogen and medroxyprogesterone or placebo CVD and MI. 138

coworkers 139 found that aspirin therapy reduced cardiovascular events by 12% in a group of 51,342 women and by 14% in a group the active treatment group. In a post hoc analysis, there was a of 44,114 men. Aspirin therapy reduced MI by 32% in men and had statistically significant 52% increase in MI in the active treatment no significant impact on stroke; it had no significant impact on MI group during the first year. 146 In the follow-up HERS II trial, more in women but reduced stroke by 17%. The decrease in stroke in than 90% of the original HERS participants were observed for 3 women was due to reductions in ischemic stroke out of proportion additional years; no evidence of cardioprotective benefit was seen to a small increase in hemorrhagic stroke. There was no significant with longer duration of follow-up. 147 effect on CVD mortality for either sex. Whereas the study found that aspirin therapy was effective in lowering the occurrence of hormone replacement formulations and delivery routes also found cardio vascular events, the differences found between men and no significant difference in IHD. 146,148-153 The Estrogen Replacement women suggest differences in mechanisms of benefit.

evaluated in the Women's Health Study 140 with the same dosing underwent coronary angiography at entry and exit. All participants strategy as in the Physicians' Health Study (100 mg every other were randomized to conjugated equine estrogen and day), which had shown a striking reduction in the risk of index MI medroxyprogesterone or placebo and observed for an average of among 22,071 healthy men. 141 In the Women's Health Study (n = 3.2 years. The results revealed no change in luminal diameter and 39,876 women aged 45 years or older), aspirin lowered the risk of no change in atherosclerosis rhotic lesion extent between the two stroke by 17%, including a 24% reduction in the risk of ischemic groups. 148 stroke with a nonsignificant increase in the risk of hemorrhagic stroke, with no reduction in fatal or nonfatal MI except for those was a double-blind, placebo-controlled trial in 199 women with no older than 65 years. Importantly, there was a significant increase IHD but elevated LDL-C (> 130 mg/dL) to determine if estrogen (40%) in gastrointestinal bleeding. Thus, a woman who is at risk is reduced the progression of subclinical atherosclerosis by use of treated differently from those at high risk with regard to platelet carotid intima-media thickness. Women (mean age, 62 years) were inhibition. All high-risk women should be prescribed a daily randomized to 17 \$ -estradiol aspirin (75 to 325 mg) unless it is contraindicated. If the woman is aspirin intolerant, clopidogrel should be used (see Table 24-3). Women classified as at risk who are younger than 65 years should not take aspirin for primary MI prevention but may consider it when the benefit of stroke prevention outweighs the increased bleeding risk associated with aspirin. In women at risk aged 65 years or older, daily aspirin reduces MI and stroke significantly;

inflammation-mediated autoimmune diseases, such as rheumatoid however, there was an almost comparable occurrence of inflammation in IHD in women. The relative risk of IHD events preventing ischemic stroke is preferred in at-risk women of all ages increases proportionally with rising levels of hsCRP and acts because aspirin has no effect on cardiovascular mortality. 139 synergistically with other risk factors to accelerate IHD risk in Aspirin resistance was found to be four times more prevalent in women. 120,124-128 A number of inflammatory markers, including women than in men in a study by Dorsch and coworkers, 142

The US Preventive Services Task Force (USPSTF) recommends 24 The use of multiple biomarkers appears to improve IHD risk the use of aspirin for women aged 55 to 79 years when the potential benefit of a reduction in ischemic stroke out weighs the potential harm of an increase in gastrointestinal hemorrhage. 143 The USPSTF concluded that the current evidence is insufficient to assess the balance of benefits and harms of aspirin for CVD prevention in men and women 80 years or older. The USPSTF recommends against the use of aspirin for stroke prevention in women younger than 55 years and for MI prevention in men younger than 45 years.

Previously, there were few data on the use of aspirin for primary addition, no difference was observed between the observational on extrapolation from large studies primarily done in men progesterone. 145 In 1998, the results from the Heart and demonstrating a reduction in risk for MI but not stroke in men of Estrogen/Progestin Replacement Study (HERS), a secondary middle age or older at intermediate or high risk. Investigators in prevention randomized trial designed to address whether estrogen the Nurses' Health Study observed a decreased risk of first MI and progestin would reduce IHD events in postmenopausal death in women who took one to six aspirins per week. 137 In the women with known CHD, called the role of hormone therapy into or placebo, in addition to hypertension treatment. The HOT study and prospectively observed for the primary outcome of nonfatal MI showed an association between aspirin use and reductions in major and death. The average age of the study participants was 67 years; 19% were diabetic, and 13% were current smokers. During a follow-A sex-specific meta-analysis conducted by Berger and up period of 4.1 years, no group outcome differences were found, despite a decrease in TC and LDL-C and an increase in HDL-C in

Subsequent to the HERS trial, many other trials with different and Atherosclerosis (ERA) trial was an invasive angiographic The use of aspirin for primary prevention in women was study in which 309 women with CAD (defined as stenosis > 30%)

The Estrogen and Prevention of Atherosclerosis Trial (EPAT) 154

160 mg/dL and observed for 2 years. Results demonstrated that the in observational studies, the age range at enrollment was 30 to 55 estradiol treatment group had greater carotid intima-media years. Finally, the average BMI was 29 in the WHI, with nearly thickness regression compared with placebo (P = 0.046), and among one third being obese, placing these women at relatively higher those not receiving statin medication, the difference was even IHD risk compared with those in the Nurses' Health study, in larger (P = 0.002). No difference was seen between estradiol and which the average BMI was 25. All of these factors may have placebo groups for the women receiving lipid-lowering therapy. contributed to variable findings between the trials and These findings suggest that among younger menopausal women observational reports. without established atherosclerosis, estradiol may be of benefit, although of a lesser magnitude compared with lipid-lowering 24 therapy.

randomized double-blind placebo-controlled trial of more than starting hormone therapy closer to menopause elicits a 27,000 healthy women aged 50 to 79 years, assessed the risks and cardioprotective effect. In the Early versus Late Inter- Ş vention benefits of hormone therapy with respect to CVD, stroke, breast Trial and colorectal cancer, and osteoporotic fractures. 155,156 There were two parallel clinical trials: estrogen-progestin and estrogen-only ° compared with placebo, based on a woman's hysterectomy status. years (the early cohort) or more than 10 years since menopause A total of 16,608 women were randomized to receive conjugated equine estrogen 0.625 mg plus medroxyprogesterone 2.5 mg or placebo, and 10,739 women with a prior hysterectomy were randomized to receive conjugated equine estrogen 0.625 mg or Kronos Early Estrogen Prevention Study (KEEPS) is a multiplacebo in the estrogen-only study. The primary outcome of the center, randomized study during a 5-year period in which 720 trial was nonfatal MI or CHD death. The duration of the follow-up women within 3 years of their final menstrual period will be for WHI was planned for 8.5 years; however, the estrogen-randomized to pill and transdermal patch hormone therapy progestin study was stopped after 5.2 years because of a global versus placebo; it also uses carotid intima-media thickness as an adverse event outcome that included an increased incidence of outcome in addition to other variables. 161 breast cancer. For IHD events, there was no significant difference in adjusted nonfatal MI and IHD deaths between the randomized groups (HR, 1.24; 95% CI, 0.97-1.60). 155 The estrogen-alone trial continued but was also subsequently stopped early after 6.8 years of follow-up because of an increased risk of stroke. In the estrogenalone trial, there was no increase in IHD risk (HR, 0.91; 95% CI, 0.75-3)

A post hoc analysis of WHI revealed fewer IHD events when postmenopausal hormone therapy was initiated closer to menopause. 157 Although this analysis was not statistically significant, it was not prospectively designed or powered to test this question. Nevertheless, the use of hormones closer to the time of menopause (<10 years) evidenced a lower risk of IHD compared with those who began therapy farther from menopause (>20 years) for both the estrogen-only and estrogen-progestin trials. The results of this analysis have provided the rationale for what has come to be termed the timing hypothesis (the timing of hormone therapy initiation closer to menopause may be cardioprotective, whereas initiation later may be adverse).

In an additional substudy within the WHI, 1064 women from the estrogen-only trial aged 50 to 59 years at study enrollment underwent coronary artery calcium scans. 158 The average age of the women was 64 years, and the mean duration of follow-up was 7.4 vears. Women who had been randomized to conjugated equine estrogen 0.625 mg had a significantly lower mean coronary calcium score of 83.1 compared with those taking placebo, 123.1 (P = 0.02). Women with at least 80% medication compliance were found to have 60% lower odds of more extensive coronary artery calcium (P = 0.004). These results are also supportive of the hypothesis that hormone therapy initiated during the menopause transition (ie, between 50 and 59 years old) may play a role in reducing the atherosclerotic plaque burden, although this hypothesis has not yet been directly tested in humans.

There are differences between the WHI clinical trial and observational reports that may account for variability between the study results. Patients with menopausal symptoms were excluded in the WHI, whereas in observational studies such as the Nurses' Health Study, they were predominant. In the WHI, only 16% of study participants were within 5 years of their last menses, with the average being 12 years, compared

with the observational studies in which enrollment was 407 largely within 5 years of menses. 159 Another way to examine these data is to evaluate differences in average age between the trials

(1 mg/day) or placebo plus lipid-lowering therapy with LDL-C > and observational data. The average age in the WHI was 63 years;

Two clinical trials are underway to address the timing of The Women's Health Initiative (WHI), a prospective - hormone therapy initiation. Both studies are addressing whether with Estradiol (ELITE), 504 women within

> (the late cohort) will be randomized to receive estradiol or placebo. In the 3-year follow-up period, outcome measures of 3this trial will include carotid intima-media thickness. 160 The

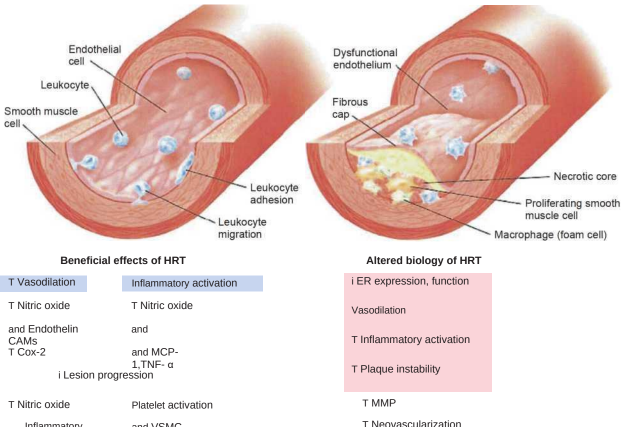
Reproductive Hormones and Premenopausal Women

Estrogen administration, in animal models, directly prevents atherosclerosis. 162 In the cardiovascular system, when estrogen receptors are activated on endothelial and myocardial cells, it has been shown to have antioxidant effects and improved endothelial cell injury recovery. 162 In specific pathways, these receptors modulate a rapid vasodilatory response through nitric oxide and also have long-term effects by increasing endothelial cell growth and inhibiting smooth muscle cell proliferation. Estrogen reduces LDL-C oxidation and binding, reduces platelet aggregation, and increases cyclooxygenase 2 activity. 162 There is relatively less known about cardiovascular actions of progesterone and progestins, and there appear to be differential effects depending on whether pre-existing atherosclerosis is present (Fig. 24-3).

IHD risk is elevated in premenopausal women with the disruption of ovulatory cycling, indicated by estrogen deficiency and hypothalamic dysfunction, 6 or irregular menstrual cycling. 163 Animal studies also demonstrated that relative estrogen deficiency is associated with atherosclerosis. These studies demonstrated that hypoestrogenemia in female monkeys is associated with loss of normal coronary artery dilation or even constriction in response to an endothelial stimulus. 164 · 166

In the WISE study, 95 premenopausal women were evaluated for hypothalamic hypoestrogenemia, defined as estradiol < 184 pmol/L (50 pg/mL), follicle-stimulating hormone < 10 IU/L, and luteinizing hormone < 10 IU/L. Premenopausal women with significant angiographic CAD had significantly lower blood levels of estradiol, bioavailable estradiol, and follicle-stimulating hormone (all P < 0.05), consistent with ovulatory dysfunction, compared with women without angiographic CAD. Hypothalamic hypoestrogenemia was significantly more prevalent among the women with CAD than among those without CAD (69% versus 29%, respectively; P = 0.01). These findings suggest that low estrogen levels due to disruption of ovarian function may be an IHD risk factor for premenopausal women.

Reproductive events in a woman's life are associated with changes in metabolic and cardiovascular function, such as blood lipids, blood pressure, and blood glucose. Despite these



Inflammatory and VSMC cell adhesion proliferation

and LDL oxidation/binding

FIGURE 24-3 The beneficial and thrombogenic effects of estrogen. CAMs, cell adhesion molecules; Cox-2, cyclooxygenase 2; ER, estrogen receptor; HRT, hormone replacement therapy; LDL, low-density lipoprotein; MCP-1, monocyte chemoattractant protein 1; MMP, matrix metalloproteinase; TNF-a, tumor necrosis factor -a; VSMC, vascular smooth muscle cell. (From Ouyang P, Michos ED, Karas RH: Hormone replacement therapy and the cardiovascular system: lessons learned and unanswered questions. J Am Coll Cardiol 47:1741, 2006.)

transient changes, a large prospective study in 1987 found no important association between variables such as age at menarche, age at first birth, and parity and IHD, whereas established risk factors showed the expected relationships. 63 Another analysis of two prospective studies, however, concluded that six or more pregnancies are associated with a small but consistent increase in the risk of IHD and CVD. 167

Polycystic ovary syndrome (PCOS) is prevalent in 10% to 13% of women and is linked with a clustering of risk factors, incident type 2 diabetes mellitus, 168 and adverse IHD events in the postmenopausal period. 169 The cardiometabolic syn drome is a clustering of risk factors including at least three of the following: insulin resistance, dyslipidemia (elevated TGs, low HDL-C), hypertension, or abdominal obesity; it is frequently associated with alterations in endogenous estrogens and androgens in women. 163,170,171

In the NHLBI-sponsored WISE study of 390 postmenopausal women with features of PCOS, 104 had clinical features of PCOS as defined by biochemical evidence of hyperandrogenemia and history of irregular menses. Cumulative 5-year cardiovascular event-free survival was 79% for PCOS women (Fig. 24-4; n = 104) versus 89% for women without PCOS (n = 286; P =0.006). PCOS women with

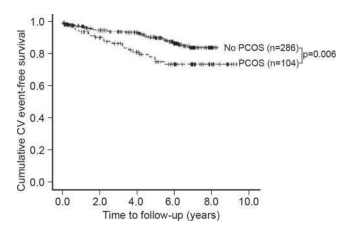


FIGURE 24-4 Cumulative unadjusted cardiovascular death or MI-free survival in postmenopausal women with or without clinical features of PCOS. (From Shaw LJ, Bairey Merz CN, Azziz R, et al: Postmenopausal women with a history of irregular menses and elevated androgen measurements at high risk for worsening cardiovascular event-free survival: results from the National Institutes of Health-National Heart, Lung, and Blood Institute sponsored Women's Ischemia Syndrome Evaluation. J Clin Endocrinol Metab 93:1276, 2008.)

elevated hsCRP had a 12.2-fold higher risk of CVD death or nonfatal MI compared with non-PCOS women and lower levels of hsCRP. PCOS status remained a significant predictor (P < 0.01) in prognostic models including diabetes, waist circumference, hypertension, and angiographic CAD as covariates. These data suggest that recognition of clinical features of PCOS in postmenopausal women may facilitate an opportunity for early risk factor intervention for the prevention of IHD.

Contraceptive Hormone Therapy

Contraceptive hormones, most commonly known as the pill, were first introduced in the 1960s to provide high hormonal blood levels that suppress ovulation, implantation, and therefore pregnancy. Contemporary oral contraceptives are considered low dose compared with first generations but remain fairly high in estrogen doses compared with menopausal hormone replacement therapy, which typically contains one tenth the dose of oral contraceptives. In addition, oralcontraceptives have a synthetic form of progestin, which ranges from the older potent androgenic progestins to newer generation progestins that are aldosterone antagonists with anti-androgenic and diuretic properties. ¹⁷² The hormones in oral contraceptives affect the cardiovascular system indirectly through their impact on CAD risk factors, such as the lipid profile, blood pressure, thrombosis, vasomotion, and arrhythmogenesis.

Lipid Effects

Changes and the amount of alteration in the lipid profile are dose dependent but also vary according to the delivery route. For example, transdermal contraceptive hormone delivery is relatively less potent than oral because it bypasses first metabolism in the liver. ¹⁶² Oral contraceptives cause hepatic apolipoprotein upregulation through the genomic pathway, which alters the lipid profiles. ^{162,173,174} Estrogen passes through the lipid membrane and binds receptors located in the nucleus, which either activates or suppresses gene transcription . Lowdose oral contraceptives of combined estrogen progestin formulations have demonstrated reductions in HDL-C and small increases in LDL-C and TGs compared with higher dose oral contraceptives. ^{175,176}

Blood Pressure

Normotensive women taking oral contraceptives have an increase in blood pressure with use up to 7 to 8 mm Hg. ^{177,178} More recently, the use of oral contraceptives with newer generation progestins (drospirenone) with antimineralocorticoid diuretic effect has been associated with lower blood pressure (a decrease ranging up to 4 mm Hg in systolic blood pressure) and lower body weight. ¹⁷⁹⁻¹⁸¹ Whereas these results are suggestive of a possible cardioprotective effect, more work is needed.

Thrombosis

Oral contraceptives have an increased risk of venous -thromboembolism and prothrombotic risk through the mechanism of increasing prothrombin and decreasing antithrombin III. ¹⁸² Compared with nonusers, users of oral contraceptive formulations with less than 50 | ig estrogen have a fourfold higher risk of venous thromboembolism (95% CI, 2.77-4.00). ¹⁸³ Third-generation (desogestrel or gestodene) progestins are twice as likely to provoke a nonfatal venous thromboembolism compared with second-generation (levonorgestrel) progestins, adjusting for smoking and BMI. ¹⁸⁴

Coronary Vasomotion

Cyclical circulating levels of endogenous and exogenous estrogen have been associated with migraine headaches, Raynaud phenomenon, and Prinzmetal angina. ^{185,186} Animal and human studies demonstrated that low endogenous estrogen levels can exacerbate endothelial dysfunction, and exogenous estrogen replacement has been shown to eliminate this effect. ^{166,187-189} Primate studies have also shown an adverse coronary

vasoconstrictive effect with medroxyprogesterone that was not apparent with progesterone. ^{190,191} Long-term studies in humans are needed because it is unknown if reproductive hormones play a role in maintaining or improving coronary or peripheral endothelial function in humans.

Arrhythmogenesis

Drug-induced QT interval prolongation and drug-induced arrhythmias are relatively more prevalent in women and contribute to a lifelong higher risk of sudden cardiac death associated with electrocardiographic QT prolongation compared with men. ¹⁹² Although data are conflicting, hormone therapy with estrogen alone usually produces a prolongation of the QT interval, and estrogen plus progesterone has no significant effects on the QT interval but reduces QT dispersion. ^{193,194} To date, studies have not been directed at the impact of oral contraceptives and mechanisms of arrhythmogenesis.

Figure 24-5 depicts the known mechanisms whereby contraceptive hormones affect the cardiovascular system, including effects on atherosclerosis, thrombosis, vasomotion, and arrhythmogenesis. ¹⁸¹ Table 24-4 lists the prescribing guidelines for hormonal contraceptives in women with elevated cardiovascular risk.

Selective Estrogen Receptor Modulators

Depending on the tissue, selective estrogen receptor modulators (SERMs) have both estrogen agonist and antagonist properties. The SERMs available in the United States and Canada are raloxifene used for prevention and treatment of osteoporosis and breast cancer prevention, tamoxifen used to prevent recurrence of estrogen receptor-positive breast cancer, and toremifene used for advanced breast cancer.

SERMs have been found to have favorable changes in regard to the lipid profiles. Raloxifene lowers TC and LDL-C; however, it does not change HDL-C or TGs. ¹⁹⁵ In addition, raloxifene does not raise hsCRP, in contrast to hormone replacement therapy; raloxifene also reduces homocysteine levels. ¹⁹⁶

Raloxifene is the most studied SERM with respect to the cardiovascular system. The Multiple Outcomes of Raloxifene Evaluation (MORE) trial was a 4-year follow-up study designed to determine the effect of raloxifene versus placebo on bone density in postmenopausal women with osteoporosis . Although this trial was designed to evaluate fracture risk, the incidence of major cardiovascular events (MI, coronary bypass surgery, percutaneous angioplasty, or stroke) was also reported. No significant differences in these events were observed between raloxifene doses (60 mg or 120 mg) and placebo. ¹⁹⁵ In a subset of women with higher risk defined as prior cardiovascular event, revascularization procedure, or multiple risk factors, raloxifene was found to have a 40% significantly reduced event rate compared with placebo. ¹⁹⁵

A subset of the MORE trial was observed for an additional 4 years in the Continuing Outcomes Relevant to Evista (CORE) trial to evaluate raloxifene and invasive breast cancer risk. ^{197,198} The combined 8-year incidence of major cardiovascular events was similar between the raloxifene group and placebo (5.5% versus 4.7%, respectively). In addition, no difference was found between treatment and placebo when IHD and stroke events were analyzed separately.

The Raloxifene Use for The Heart (RUTH) trial was the first trial designed to evaluate raloxifene and the effect on cardiovascular events as a primary outcome. ¹⁹⁹ This trial enrolled 10,101 postmenopausal women with known CHD or



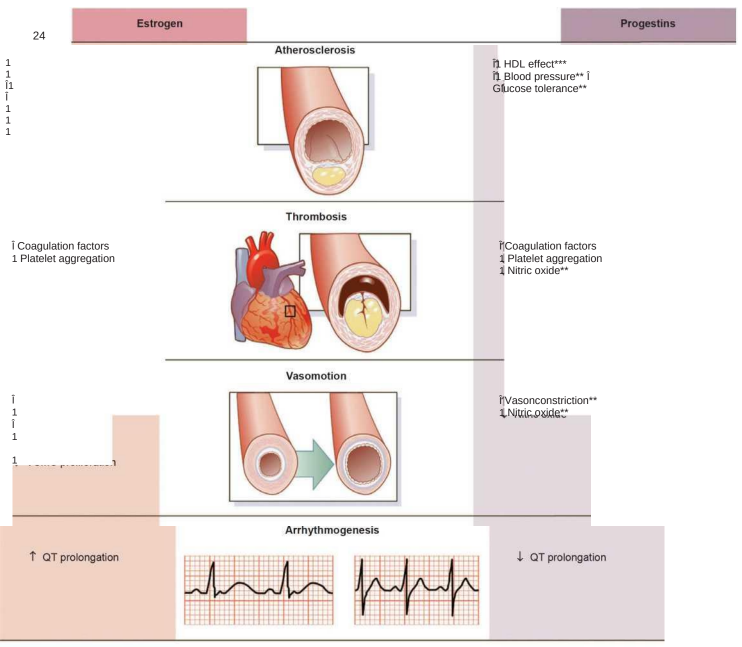


FIGURE 24-5 Impact of hormonal contraception on mechanisms of cardiovascular disease. Cox-2, cyclooxygenase 2; HDL, high-density lipoprotein; LDL, low-density lipoprotein; VSMC, vascular smooth muscle cell. *Dependent on delivery route of estrogen. **Dependent on type of progestin. ***Dependent on the dose of estrogen. (From Shufelt CL, Bairey Merz CN: Contraceptive hormone use and cardiovascular disease. J Am Coll Cardiol 53:221, 2009.)

LDL oxidation
LDL binding
Lipoprotein*,***
Blood pressure
Oxidation damage
VSMC proliferation
Glucose tolerance***
Nitric oxide
Endothelin
Cox-2
Neuroendocrine
response
VSMC proliferation

TABLE 24-4	Summary of Hormonal Contraceptive Prescribing Guidelines for Women with Elevated Cardiovascular Risk
Hypertension	Well-controlled blood pressure in women < 35 years of age and otherwise healthy, nonsmoking trial of oral contraceptives. Monitor blood pressure and if controlled after starting, oral contraceptives may be continued. If blood pressure is not well controlled, alternative methods such as progestin-only pills or intrauterine device (IUD) may be started.
Dyslipidemia	LDL-C > 160 mg/dL or multiple cardiac risk factors alternative nonhormonal contraceptive methods, such as an IUD.
Diabetes	Diabetes type 1 or 2, oral contraceptives are appropriate for use <i>only</i> in otherwise healthy nonsmokers < 35 years of age. Otherwise, progestin-only or IUD may be started.
Tuxedo	Smoking and > 35 years of age alternative nonhormonal contraceptive methods, such as an IUD. Smokers < 35 years of age are not addressed.
Obesity	Obesity (BMI > 30 kg/m ·) alternatives nonhormonal contraceptive methods such as progestinonly contraception or IUD. Obesity is thought to be an independent risk factor for venous thromboembolism.
Women over 35 years old	Healthy, nonsmoking women oral contraceptives with < 50 ^ g ethinyl estradiol remain safer than pregnancy and can be continued until 50 to 55 years of age or until menopause after reviewing risks and benefits.

From Shufelt CL, Bairey Merz CN: Contraceptive hormone use and cardiovascular disease. J Am Coll Cardiol 53:221, 2009.

risk factors for a median follow-up of 5.6 years and found that raloxifene did not reduce cardiovascular event risk (death from coronary causes, nonfatal MI, or hospitalization from acute MI). 200 In a post hoc analysis, women < 60 years old had significantly fewer cardiovascular events when randomized to raloxifene versus placebo (HR, 0.59; 95% CI, 0.41 to 0.83; P = 0.003). 200 These findings suggest, similar to the animal, observational, and clinical trial data, that there may be a CVD benefit with hormonal therapy at younger ages.

GUIDELINES FOR PREVENTION OF CARDIOVASCULAR DISEASES IN WOMEN

The 2007 AHA update recommends an integrated scheme for a general risk stratification approach to the female patient that classifies her as at high risk, at risk, or at optimal risk, focused on the high average lifetime risk for CVD, approaching one in two women. The guidelines recommend aggressive risk factor modification through lifestyle modification and phar macotherapy consistent with the ATP III guidelines, including smoking cessation, exercise, weight reduction, and a diet rich in fruits and vegetables. The guidelines also emphasize the importance of aggressive control of blood pressure and lipids through lifestyle modification and pharmacotherapy. Aspirin therapy is recommended in high-risk women if there are no contraindications. Aspirin therapy is also recommended for women > 65 years, if blood pressure is controlled and benefit for ischemic stroke and MI prevention is likely to outweigh the risk of gastrointestinal bleeding and haemorrhagic stroke, and for women < 65 years when benefit for ischemic stroke prevention is likely to outweigh adverse

TABLE 24—5 Class III Interventions (Not Useful/Effective and May Be Harmful) for Cardiovascular Disease or Myocardial Infarction Prevention in Women

Menopausal Therapy

Hormone therapy and selective estrogen receptor modulators (SERMs) should not be used for the primary or secondary prevention of CVD (Class III, Level A).

Antioxidant Supplements

Antioxidant vitamin supplements (eg, vitamin E, vitamin C, and beta carotene) should not be used for the primary or secondary prevention of CVD (Class III, Level A).

Folic Acid*

Folic acid, with or without B $_{6}$ and B $_{12}$ supplementation, should not be used for the primary or secondary prevention of CVD (Class III, Level A) .

Aspirin for Myocardial Infarction in Women < 65 Years of Age A

Routine use of aspirin in healthy women < 65 years of age is not recommended to prevent myocardial infarction (Class III, Level B).

- *Folic acid supplementation should be used in the childbearing years to prevent neural tube defects.
- Tor recommendation for aspirin to prevent CVD in women >65 years of age or stroke in women <65 years of age. see Table 24-3.
- From Mosca L, Banka CL, Benjamin EJ, et al: Evidence-based guidelines for cardiovascular disease prevention in women: 2007 update. *Circulation* 115:1481, 2007

effects of therapy. Beta blockers are recommended in women after MI, acute coronary syndrome, or left ventricular dysfunction with or without heart failure symptoms. Similarly, angiotensin-converting enzyme inhibitors are recommended in women after MI and in those with heart failure, left ven tricular dysfunction, or diabetes mellitus. Hormone therapy, antioxidants, and vitamins should not be used for primary or secondary prevention in women (Table 24-5).

CONCLUSION

CVD is the leading cause of death among women in the United States, and women receive fewer preventive recommendations, such as lipid-lowering therapy, aspirin, and lifestyle advice, than do men with similar Framingham risk scores. Both the Framingham risk score and the Reynolds score can be used to estimate CVD risk in women. Treatment of dyslipidemia, hypertension, and diabetes is similar between women and men. The use of aspirin for women aged 55 to 79 years is recommended when the potential benefit of a reduction in ischemic strokes outweighs the potential harm of an increase in gastrointestinal hemorrhage. Lifestyle is an important component of prevention of IHD; women should accumulate a minimum of 30 minutes of moderate-intensity physical activity (eg, brisk walking) daily, consume a heart-healthy diet, and avoid smoking or passive smoke. Hormone therapy used for contraception, management of menopause symptoms, or other clinically indicated conditions should not be used for CVD prevention. Sex-specific guidelines exist for women for the prevention of CVD and should be used by health care providers.

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CHAPTER 25

Cardiovascular Aging: The Next **Frontier in Cardiovascular Prevention**

KEY POINTS Samer S. Najjar, Edward G. Lakatta, and Gary Gerstenblith The incidence and prevalence of

- most cardiovascular diseases increase with advancing age.
- Age is the dominant risk factor for cardiovascular diseases, but age has traditionally been viewed as a nonmodifiable risk factor.
- Physiologic aging of the cardiovascular system is associated with myriad structural and functional alterations in the heart and in the blood vessels, which are themselves risk factors for cardiovascular diseases and thus may help explain the "risky" aspects of aging.
- The hallmarks of arterial aging include thickening and stiffening of the walls of the central arteries and endothelial dysfunction.
- Cardiac aging includes structural alterations in the heart, deficits in reserve functions, alterations in calcium regulation, and increased generation of reactive oxygen species during stress.
- Clinical interventions to retard cardiovascular aging include lifestyle changes (e.g., exercise, smoking cessation) and pharmacologic interventions (e.g., renin-angiotensinaldosterone antagonists, antihypertensive medications, statins).

 Future studies should examine whether interventions aimed at retarding cardiovascular aging can have a positive impact on the adverse cardiovascular effects of accelerated cardiovascular aging and attenuate the impact

The epidemic cardiovascular diseases has taken on a global dimension and is no longer restricted Western societies. Car diovascular diseases now represent more than 30% of all deaths worldwide, and by the year 2020, they are expected to surpass infectious diseases as the leading cause of mortality disability. According to the World Health Report, cardiovascular diseases were responsible for 15 million annual deaths worldwide, of which 9 million were

developing countries and 2 million economies in transition. This situation is expected to worsen as the world population in both industrialized and developing countries is aging. For example, in the United States, 35 million people are older than 65 years, and the number of older Americans is expected to double by the year 2030. In many developing countries, the older population progressively becoming more pre dominant as life expectancy increases.

The clinical and economic implications of this demographic shift are staggering because age is the most potent individual risk factor for cardiovascular diseases, especially

individuals older than 50 years. ² Both the incidence and prevalence of hypertension, coronary artery disease, congestive heart failure, and stroke increase exponentially with age. In older community dwelling healthy volunteers, the incidence of silent coronary atherosclerosis. assessed by combined electrocardiographic treadmill stress testing and thallium perfusion imaging, increases dramatically with age. 3

Age influences not only the incidence and prevalence of coronary atherosclerosis but also the severity and prognosis of this disease. In survivors of a myocardial infarction, age is an independent predictor of

age as the dominant risk factor for cardiovascular diseases. Importantly, this should help change our view of aging from an immutable risk factor to one that is amenable to modification and prevention.

short- and long-term morbidity, mortality, and disability, even after adjustment for the infarct size and location, the number of dis eased vessels, and the extent of coronary artery disease. ⁴Similar considerations pertain to the impact of age on other cardio vascular diseases, such as hypertension, congestive heart failure, and stroke.

In spite of the fact that age is the dominant risk factor for cardiovascular diseases, most of the research efforts on prevention of these diseases have ignored age and have focused instead on the development of interventions that target "traditional" cardiovascular risk factors (such as hypertension and hyperlipidemia) or identification of newer risk markers. This is because age has usually been

viewed as an unmodifiable risk factor and therefore one that is not amenable to prevention or treatment. However, this concept has recently been challenged by an emerging school of thought, which proposes that aging can indeed be construed as a modifiable risk factor. ⁵

The key paradigm shift is that the components of aging associated with cardiovascular disease risk should no longer be simplistically attributed to an increased time of exposure to other established cardiovascular risk factors. Instead, the risky components of aging need to be recognized as accelerated or dysregulated age-associated alterations in the cardiovascular system, at the molecular, enzymatic, bio chemical, cellular, histologic, and organis mal levels. In this chapter, we review some of these alterations, at both the microscopic and clinical levels, with a view towards illustrating the overlapping features of aging with some cardiovascular diseases and risk factors. We provide an integrated

416 view of the risky components of cardiovascular aging and of agedisease interactions, which are in dire need of targeted preventive and therapeutic strategies.

EPIDEMIOLOGY OF CARDIOVASCULAR DISEASES IN OLDER INDIVIDUALS

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The incidence and prevalence of most cardiovascular diseases increase with advancing age in both sexes. ⁶ For example, the incidence of heart failure in the Framingham Heart Study (1980-2003) per 1000 person-years was 9.2 and 4.7 in men and women between the ages of 65 and 74 years, 22.3 and 14.8 in men and women between the ages of 75 and 84 years, and 41.9 and 32.7 in men and women older than 85 years. The prevalence of heart failure according to NHANES (2003 2006) was 2% and 1% in men and women between the ages of 40 and 59 years, 9% and 5% in men and women between the ages of 60 and 79 years, and 15% and 12% in men and women older than 80 years.

The prevalence of hypertension (defined as systolic blood pressure higher than 140 mm Hg or diastolic blood pressure higher than 90 mm Hg) in NHANES (2003-2006) was 39% and 38% in men and women between the ages of 45 and 54 years, about 54% in men and women between the ages of 55 and 64 years, 65% and 71% in men and women between the ages of 65 and 74 years, and 65% and 77% in men and women older than 75 years.

The incidence of stroke in ARIC (1987-2001) per 1000 personyears was 2.4 for white men and women between the ages of 45 and 54 years and 9.7 and 7.2 for black men and women in the same age range. These numbers increased to 6.1 and 4.8 for white men and women between the ages of 55 and 64 years and 13.1 and 10.0 for black men and women in this age range. The incidence of stroke increased to 12.1 and 9.8 in white men and women between the ages of 65 and 74 years and to 16.2 and 15 in black men and women in this age range. The prevalence of stroke in NHANES (2003-2006) was 1% and 6% in men and women between the ages of 40 and 59 years, 7% in men and women between the ages of 60 and 79 years, and 15% and 13% in men and women older than 80 years.

Last, the prevalence of coronary heart disease in NHANES (2003-2006) was 7% in both men and women between the ages of 40 and 59 years, 25% and 17% in men and women between the ages of 60 and 79 years, and 37% and 23% in men and women older than 80 years.

Importantly, age also influences the prognosis after a cardiovascular event, such that within 1 year after a myocardial infarction, the rate of death is 8% and 12% in white men and women between the ages of 40 and 69 years and 14% and 11% in black men and women in this age range. This rate increases to 27% and 32% of white men and women older than 70 years and to 26% and 28% of black men and women in this age range. Similarly, the risk of heart failure 5 years after a myocardial infarction increases with age. It is 7% and 12% in men and women between the ages of 40 and 69 years and 22% and 25% in men and women older than 70 years.

AGING OF CENTRAL ARTERIES IN APPARENTLY HEALTHY HUMANS

The large central arteries had originally been constructed as pipes that simply allowed blood to flow from the heart to distal organs and thus were originally dubbed "conduit" arteries. Changes in their mechanical properties with aging were considered to result largely from passive wear and tear in response to the unrelenting stress and strain incurred with each heartbeat throughout life. However, recent insights into

	Humans > 65	4g/ng Monkeys 15-20 years	Rats 24-30 months	Rabbits 2-6 years
Luminal dilation	+	+	+	+
T Stiffness	+	+	+	+
Endothelial dysfunction	+	+	+	+
Diffuse intimately thickening	+	+	+	+
Collagen degradation	+	+	+	+
Collagen deposits	on +	+	+	+
VSMC number	+	+	+	+
Macrophages	+	-	-	-
T cells	+	-	-	-
T Local angiotensin II-ACE	+	+	+	+
MMP dysregulation	+	+	+	?
T MCP-1/CCR2	+	+	+	+
T ICAM	+	+	+	?
T TGF- β	+	+	+	+
T NADPH oxidase	+	+	+	+
F VEGF	+	?	?	+
F Nitric oxide bioavailability	+	+	+	+

information unknown. ACE, angiotensin-converting enzyme; ICAM, intercellular adhesion molecules; MCP-1, monocyte chemotactic protein 1; MMP, matrix metalloproteinase; TGF- β , transforming growth factor - β ; VEGF, vascular endothelial growth factor; VSMC, vascular smooth muscle cell.

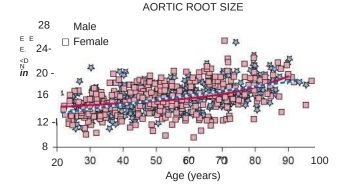
the multiple signaling mechanisms that modulate their ability to adapt, repair, and remodel and that govern their structural and functional properties have left us marveling at these dynamic organs, which play pivotal roles in regulating pulse flow and modulating vascular impedance. The age-associated changes in central arterial structure and function in primates and in apparently healthy humans are summarized in Table 25-1. ¹ Some of these features are discussed in the following sections.

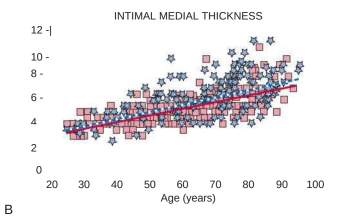
Overall cardiovascular structure and function vary -dramatically among older individuals. ⁷ Identification of cardio vascular structural and functional changes that reflect an "aging process" is a formidable task. Neither functional differences among individuals in cross-sectional studies nor changes over time within a given individual in longitudinal studies are necessarily manifestations of an aging process. Rather, interactions among aging, disease, and lifestyle must be considered in interpreting age-associated changes in cardiovascular structure and function as measured in various studies. ⁸

Lumen Diameter

Cross-sectional studies show that on average, the central aorta dilates with age, leading to an increase in lumen size ⁹ (Fig. 25-1A). Studies found an inverse and independent

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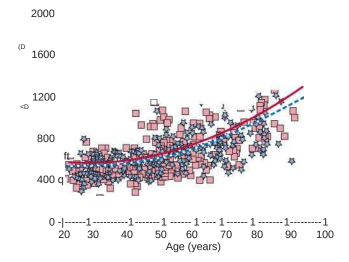


FIGURE 25-1 Age-associated changes in arterial structure and function in men (blue stars) and women (red squares) in the Baltimore Longitudinal Study of Aging. Best fit regression lines (quadratic or linear) are shown for men (solid lines) and women (dotted lines). A, Aortic root size (measured by M-mode echocardiography) indexed to body surface area. B, Common carotid intima-medial thickness (measured by B-mode ultrasonography), C. Carotid-femoral pulse wave velocity (an index of central arterial stiffness).

association between aortic root diameter and pulse pressure, suggesting that luminal size may play a role in the pathogenesis of systolic hypertension. ¹⁰ This provocative concept awaits the results of longitudinal studies to clarify the significance of the cross-sectional relationship between aortic root diameter and pulse pressure, which could have important implications for putative future therapies to delay or to prevent systolic hypertension, the dominant form of hyper tension in older individuals.

Wall Thickness

Aging is also associated with thickening of the walls of the central arteries (Fig. 25-1B). Intima-media thickness rises nearly threefold between the third and ninth decades of life, an increase that is mainly attributable to an increase in intimal thickness.¹¹ The age-associated intima-media thickening is often ascribed to atherosclerosis, and intimamedia thickness is often equated with subclinical atherosclerosis. This is because a growing body of literature has shown that traditional cardiovascular risk factors and prevalent 25 cardiovascular diseases are associated with intima-media thickening and that intima-media thickening is a potent and independent predictor of adverse cardiovascular events. However, intima-med thickness is only modestly correlated with the extent and severity af coronary artery disease, 12 and it can be influenced by factors other than atherosclerosis (e.g., aging, hypertension).

For example, intima-media thickening increases with advancing age in animal models that are devoid of atheroscle- rosis, 13 thus indicating that these age-associated alterations are due to the aging process and not to superimposed atherosclerosis. This thickening as due to myriad age-associated biochemical, cellular, and morphologic changes in the arterial wall, which are modulated by the same factors that have been implicated in the genesis of various cardiovascular dis- eases.14 Thus, întima-media thickening should not be constructed

dis- eases. 14 Thus, intima-media thickening should not be constructed as synonymous with "subclinical atherosclerosis," particularly in the absence of plaques. Nonetheless, it remains a useful marker subclinical vascular disease. 15

Wall Remodeling

Although we have discussed arterial wall thickness and luminal diameter separately, arteries undergo a remodeling process, in response to hemodynamic and metabolic stimuli, that can be better characterized by an index that combines the diameter, wall thickness, and vascular mass variables than by each of these thickness, and vascular mass variables than by each of these variables individually. For example, in the carotid artery, three remodeling patterns that deviate from normal can be described16: (1) concentric remodeling, due to a smaller lumen in the setting of a normal vascular mass; (2) vascular hypertrophy, due to an increase in vascular mass, which results from wall thickening, in the setting of a normal lumen size; and (3) eccentric hypertrophy, due to an increase in vascular mass in the setting of a dilated lumen. In a study of normotensive and untreated hypertensive Taiwanese subjects, these geometric patterns yielded prognostic information, with the two hypertrophic groups being associated with an increased risk of cardiovascular events.¹⁷ This underscores the importance of integrating the markers of arterial aging to better appreciate their clinical utility.

Arterial Stiffness

One of the hallmarks of central arterial aging is an age-associated increase in stiffness (Fig. 25-1C). Central arterial stiffness can be assessed by several invasive and noninvasive indexes. Recently, carotid-femoral pulse wave velocity has been anointed the "gold standard" for the noninvasive assessment of central arterial stiffness. 18 Pulse wave velocity has been shown to be an independent predictor of morbidity and mortality in healthy subjects and in individuals with various levels of cardiovascular risk. It is likely that arterial stiffness is not only a risk marker for cardiovascular diseases but also a risk factor for these diseases. Indeed, increased central arterial stiffness leads to an increase in the load (afterload) on the left ventricle, leading to increased hypertrophy and increased oxygen consumption. Furthermore, increased stiffness is associated with a decrease in diastolic blood pressure, which

could compromise coronary blood flow, which occurs predominantly in diastole.

The age-associated increase in stiffness has traditionally been attributed to the fraying and breakdown of elastin due to the lifelong repeated cycles of distention and recoil of the central aorta as well as the increased deposition and covalent cross-linking of collagen molecules. It is now recognized that arterial stiffening can be modulated by several factors including lifestyle considerations (eg, salt intake, exercise, weight loss), ¹⁹ signaling pathways (eg, nitric oxide), 20 inflammation, and genetics. 21 25 Interventions to prevent or to delay arterial stiffening have predominantly focused on pharmacological antihypertensive therapies. However, these strategies are aimed at lowering blood pressure, whereby the reduction in stiffness is a secondary effect due to reverse remodeling of the arterial wall in response to the lower pressures.

Because arterial stiffness is a potent predictor of mortality and morbidity, independent of blood pressure, a more direct approach that would target the stiffening process is desirable. However, there is a dearth of interventions at present that have attempted this, with the one notable exception being the advanced glycation end products cross-link breaker Alagebrium (ALT-711). This compound cleaves the covalent cross link bonds that form between adjacent collagen fibrils in the medial layer of the arterial wall and that contribute to increasing the tensile strength of the collagen peripheral blood pressures. 28 molecules. Alagebrium has been shown to reduce arterial stiffness in rodents and in nonhuman primates.

In humans with systolic hypertension and increased arterial stiffness, a randomized, double-blind, placebo-controlled study showed that treatment with Alagebrium for 8 weeks resulted in a significant decrease in arterial stiffness, without an apparent disproportionate decline in mean arterial pressure, systemic vascular resistance, heart rate, or cardiac output. 22 Although a multicenter, randomized, double-blind, placebo-controlled clinical trial failed to show an effect of Alagebrium on blood pressure, ²³ the primary effect of this compound should be a reduction in stiffness, and a reduction in stiffness may not necessarily translate into a reduction in blood pressure (at least in the short or intermediate term) because the lowering in stiffness could be accompanied by an increase in stroke volume (lower afterload on the heart), which could potentially offset the effects of a lower stiffness on blood pressure. Unfortunately, these haemodynamic features were not Arterial-Ventricular Coupling assessed in the aforementioned clinical study.

muscular arteries (eg, brachial and femoral arteries) does not (LV) structure and function. Importantly, the left ventricle and the increase with age. Thus, the manifestations of arterial aging may vary among the different vascular beds, reflecting differences in the index of this interaction, termed arterial-ventricular coupling, can structural compositions of the arteries and perhaps differences in the age-associated signaling cascades that modulate the arterial properties or differences in the response to these signals across the arterial tree.

Reflected Waves

The increased load that is imposed on the heart by increasing stiffness is due not only to an increase in characteristic impedance (intrinsic stiffness of the vessel) but also to an increase in central systolic blood pressure. The latter is due, in part, to a shift in the return of reflected waves to an earlier time during systole, which leads to an increase in central pressure augmentation and thus to a higher central systolic blood pressure and a higher central pulse pressure. It is now appreciated that the timing and amplitude of reflected waves are governed both by arterial stiffness and by properties of the distal reflecting sites (usually areas of impedance mismatch). These reflected waves play an important role in influencing the central to peripheral pressure augmentation across the arterial tree. ²⁴ Even though peripheral systolic pressure and pulse pressure increase with age, the central to peripheral pressure amplification decreases with age.

Central Blood Pressure

Recent methodological advances have made it possible to noninvasively estimate central blood pressures with relative ease. This is fueling a growing clinical interest in assessing central blood pressures. The Conduit Artery Functional Endpoint (CAFE) study showed that the combination of the calcium channel blocker amlodipine with or without the angiotensin converting enzyme (ACE) inhibitor perindopril in the treatment of hypertension achieved a greater reduction in central blood pressure than the beta blocker atenolol with or without a diuretic, even though the peripheral blood pressures were not significantly different between the two groups. ²⁵CAFE was a substudy of the Anglo-Scandinavian Cardiac Outcomes Trial (ASCOT), which showed that patients in the amlodipine-perindopril arm had better clinical outcomes than those in the atenolol-diuretic arm. ²⁶

The findings from CAFE suggest that the lower central blood pressures achieved with amlodipine-perindopril may explain, in part, the differences in outcomes between the two groups. The usefulness and feasibility of integrating measurements of central arterial blood pressures into routine medical practice are areas of intense ongoing research. As a first step, studies have established that central pressures are independent predictors of outcomes 27 and may be even better predictors of cardiovascular mortality than

Increased central arterial stiffness is associated with an increase in central systolic blood pressure and a decrease in diastolic blood pressure, resulting in a widening of the pulse pressure. This is a likely explanation of the age-associated changes in blood pressure, whereby systolic blood pressure continues to increase with advancing age, whereas diastolic blood pressure increases until the fifth decade, then levels off and starts to decrease after the age of 60 years. ²⁹ Numerous clinical and epidemiological studies in several different populations with varying prevalence of cardiovascular diseases have demonstrated that pulse pressure is an important predictor of adverse outcomes, often more potent than systolic or diastolic blood pressures. However, the clinical role of pulse pressure has remained limited to a few clinical conditions (eg, advanced heart failure, severe aortic regurgitation), although it could serve as an easily obtainable surrogate marker of stiffness.

In addition to the age-associated alterations in arterial proper ties, In contrast to the central elastic arteries, the stiffness of the advancing age is also associated with alterations in left ventricular central arteries have bidirectional constant interactions. One useful be assessed by the ratio of effective arterial elastance (Ea), a measure of the net arterial load exerted on the left ventricle, to LV end-systolic elastance (ELV), a load-independent measure of LV chamber performance. 30 Ea increases with advancing age, predominantly because of the age-associated increase in arterial stiffness, as peripheral vascular resistance and heart rate (the other main determinants of Ea) do not significantly change with age. 31

> ELV also increases with advancing age. Interestingly, even though Ea and ELV both increase with advancing age, their ratio, Ea/ELV, remains relatively unchanged across the age spectrum in men, ³² suggesting that the increases in Ea and ELV are matched. This tight coupling is thought to allow the

cardiovascular system to optimize energetic efficiency. In contrast, Plasminogen Activator Inhibitor 1 in healthy women, Ea/ELV declines slightly with advancing age, 31 which is due to a disproportionate age-associated increase in ELV compared with the age-associated increase in Ea. This suggests that in women, but not in men, aging exerts a greater impact on ventricular than arterial properties.

Therapies that improve the coupling of ventricular and vascular elastances are likely to improve cardiac function and exercise tolerance in healthy subjects. This concept is supported by two studies in healthy older subjects. Administration of the direct vasodilator sodium nitroprusside caused reductions in reflected waves (manifested as a reduction in preload), resulting in reduced cardiac volumes and higher ejection fraction at rest and during maximal exercise compared with placebo therapy. Administration of intravenous verapamil reduced noninvasive indices of arterial and ventricular systolic stiffness and improved exercise tolerance and oxygen consumption before reaching the anaerobic threshold.

Aging of Endothelial Cells

An important feature of arterial aging is age-associated endothelial cells in central arteries has not been directly assessed. advancing age. 34

Endothelial cells play a pivotal role in modulating myriad basis for cardiovascular aging in relation to thrombosis. arterial structural and functional properties, including vascular tone, vascular permeability, angiogenesis, production of extracellular matrix proteins and of growth factors, neurohormonal CARDIAC AGING signaling (nitric oxide, renin-angiotensin-aldosterone system, sympathetic nervous system), and response to inflammation. The critical role of endothelial cells in several cardiovascular diseases is increasingly being appreciated . For example, endothelial changes in the heart. dysfunction has been described in patients with hypertension and, interestingly, in the normotensive offspring of hypertensive Cardiac Aging in Humans individuals, 35 suggesting that endothelial dysfunction may With advancing age, the walls of the left ventricle increase in precede the development of clinical hypertension. Endothelial dysfunction has also been implicated in the pathogenesis of atherosclerosis and is one of the earliest pathological manifestations of this disease. Interestingly, arterial regions with low shear stress are particularly vulnerable to the development of atherosclerosis, and vascular wall shear stress is an important modulator of endothelial cell morphology and endothelial cell metabolic and synthetic functions. Endothelial dysfunction, in both the coronary and peripheral arterial beds, has been shown to be an independent predictor of future car diovascular events.

Endothelial dysfunction can also be induced by loss of telomere function. Telomeres are DNA-protein complexes that form the ends of chromosomes, and they shorten with each replicative cycle, which is why they have been proposed to be possible indicators of biological aging. Interestingly, inhibition of telomere shortening can suppress the age-associated endothelial dysfunction. ³⁶

Vascular cell senescence is increasingly recognized as an important feature of arterial aging. Thus, there is growing interest in and excitement at the promises held by regenerative therapies. Endothelial cells play a pivotal role in angiogenesis, the process through which new vessels grow from existing microvasculature. However, studies in animal models of aging indicate that angiogenesis is impaired with advancing age. Endothelial progenitor cells offer the prospects of putatively replenishing or replacing senescent endothelial cells. However, both the number ³⁷ and activity ³⁸ of endothelial progenitor cells appear to be reduced with aging, 419 suggesting an age-associated impairment in regenerative and repair capacities, which, experimentally, can be reversed with growth hormone-mediated increase in insulin-like growth factor 1 (IGF-1) levels. 39

Thrombotic cardiovascular diseases increase in incidence in the elderly, a tendency dependent on the age-related changes in vascular and hemostatic systems that include 25 platelets, coagulation, and fibrinolytic factors as well as in the endothelium. The hypercoagulability of and advanced sclerotic changes in the vascular wall may contribute to the increased incidence of thrombosis in the elderly. One of the o important key genes for aging-associated thrombosis is plas minogen activator inhibitor 1 (PAI-1), a main inhibitor of £_ fibrinolysis.

The expression of PAI-1 is not only elevated in the elderly but also significantly induced in a variety of pathological conditions associated with the process of aging. These conditions include obesity, insulin resistance, emotional stress, immune responses, and vascular sclerosis and remodeling. Several cytokines and hormones, including tumor necrosis factor- a (TNF -a), transforming growth factor- \$ (TGF -\$), -n angiotensin II, and insulin, positively regulate the gene expression of PAI-1.

The recent epidemic of obesity with aging in industrialized societies may heighten the risk for thrombotic cardiovascular disease because adipose tissue is a primary source of PAI-1 and compromise in endothelial function. In humans, the function of cytokines. Emotional or psychosocial stress and inflammation also cause the elevated expression of PAI-1 in an age-specific pattern. Nevertheless, endothelial function has been assessed in the brachial Thus, PAI-1 could play a key role in the progression of artery, where measurement of vasoreactivity by both agonist- and cardiovascular aging by promoting thrombosis and vascular flow-mediated techniques has shown that it declines with (athero)sclerosis. Further studies on the genetic mechanism of aging-associated PAI-1 induction will be necessary to define the

The preceding section focused on age-associated changes in the vasculature. Aging is also associated with structural and functional

thickness, largely because of an increase in ventricular myocyte size and an increase in vascular impedance, and this helps moderate the increase in LV wall tension. Modest increases in collagen levels also occur with aging.

A unified interpretation of identified cardiac changes that accompany advancing age in otherwise healthy persons suggests that at least in part, these are adaptive, occurring to some extent in response to arterial changes that occur with aging (Fig. 25-2). 9 Prolonged contraction of the thickened LV wall maintains a normal ejection time in the presence of the late augmentation of aortic impedance. This preserves the systolic cardiac pumping function at rest. One disadvantage of prolonged contraction is that at the time of the mitral valve opening, myocardial relaxation is relatively more incomplete in older than in younger individuals and causes the early LV filling rate to be reduced in older individuals.

Structural changes and functional heterogeneity occurring within the left ventricle with aging may also contribute to this reduction in peak LV filling rate. However, concomitant adaptations-left atrial enlargement and an enhanced atrial contribution to ventricular filling – compensate for the



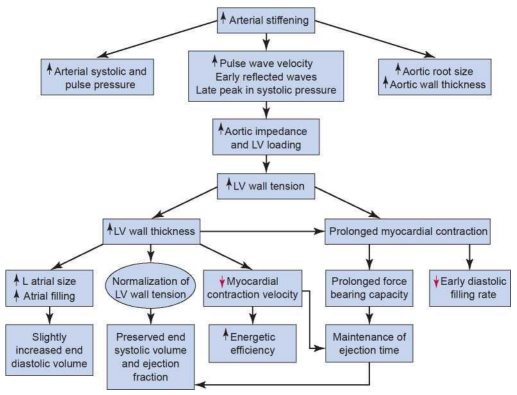


FIGURE 25-2 Arterial and cardiac changes that occur with aging in healthy humans.

vascular cells or their extracellular matrices (see later) may also hypertension that causes either systolic or diastolic heart failure). have a role in the scheme depicted in Figure 25-2.

morphometric assessments in nonfailing human hearts have shown cardiovascular system is to ensure that the heart beats faster; to extensive age-related myocyte loss and hypertrophy of the ensure that it retains a small size, by reducing the diastolic filling surviving myocytes in male hearts but preserved ventricular period, reducing LV afterload; to augment myocardial contractility myocardial mass and average cell diameter and volume in aging and relaxation; and to redistribute blood to working muscles and female hearts. These sex differences may stem, in part, from to skin to dissipate heat. Each of the deficient components of differences in the replicative potential of cardiac myocytes. cardiovascular regulation with aging, that is, heart rate (and thus of the left ventricle from patients with dilated cardiomyopathy has contractility, and redistribution of blood flow, exhibits a deficient identified more than 1800 genes displaying sexual dimorphism in sympathetic modulation. the heart. A significant number of these genes were highly represented in gene ontology pathways involved in ion transport postsynaptic beta-adrenergic signaling declines with aging. One and G protein-coupled receptor signaling. 42

Cardiovascular Reserve

Impaired heart rate acceleration and impaired augmentation of blood ejection from the left ventricle, accompanied by an acute modest increase in LV end-diastolic volume, are the most dramatic changes in cardiac reserve capacity that occur with aging in only in younger subjects (Fig. 25-3A). The heart rate reduction healthy, community-dwelling persons (Table 25-2). Mechanisms during exercise in the presence of acute beta-adrenergic blockade that underlie the age-associated reduction in maximum ejection is greater fraction are multifactorial and include a reduction in intrinsic myocardial contractility, an increase in vascular afterload, and arterial-ventricular load mismatching.

Ventricular load is the opposition to myocardial contraction and the ejection of blood; afterload is the component of load that pertains to the time after excitation, as opposed to preload, before excitation. Although these age-associated changes

reduced early filling and prevent a reduction of the end-diastolic cardiovascular reserve are insufficient to produce clinical heart volume. Age-associated changes in the tissue levels of or responses failure, they do affect its clinical presentation, that is, the threshold to growth factors (catecholamines, angioten sin II, endothelin, TGF- for symptoms and signs, or the severity and prognosis of heart S, or fibroblast growth factor) that influence myocardial or failure secondary to any level of disease burden (eg, chronic

A sizeable component of the age-associated deficit in Biological sex is a well-recognized factor in the physiology and cardiovascular reserve is composed of diminished effectiveness of pathophysiology of the cardiovascular system, including the aging the autonomic modulation of heart rate, LV contractility , and heart (reviewed in references 40 and 41). Postmortem - arterial afterload. The essence of sympathetic modulation of the Analysis of gene expression differences by sex and age in samples filling time), afterload (both cardiac and vascular), myocardial

Multiple lines of evidence support the idea that the efficiency of line of evidence stems from the observation that cardiovascular responses to beta-adrenergic agonist infusions at rest decrease with age. A second type of evidence for a diminished efficacy of postsynaptic beta-adrenergic receptor signaling is that acute betaadrenergic receptor blockade changes the exercise hemodynamic profile of younger persons to make it resemble that of older individuals. Significant beta blockade-induced LV dilation occurs

TABLE 25-2

Exhaustive Upright Exercise: Changes in Aerobic Capacity and Cardiac Regulation Between the Ages of 20 and 80 Years in Healthy Men and Women

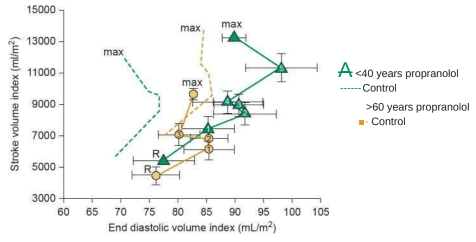
Oxygen consumption	F (50%)
(AV)O 2	F (25%)
Cardiac index	F (25%)
Heart rate	F (25%)
Stroke volume Preload EDV	No change
Afterload	T (30%) T
Vascular (PVR) Cardiac (ESV)	T (30%) T (275%)
Cardiac (EDV) Contractility	T (30%) F (60%)
Ejection fraction	F (15%)
Plasma catecholamines	Т
Cardiac and vascular responses to beta-adrenergic stimulation	F

EDV, end-diastolic volume; ESV, end-systolic volume; PVR, peripheral vascular resistance.

younger versus older subjects (Fig. 25-3B), as are the age- **421** associated deficits in LV early diastolic filling rate, both at rest and during exercise (Fig. 25-3C). It has also been observed in older dogs that the age-associated increase in aortic impedance during exercise is abolished by acute beta-adrenergic blockade.

Apparent deficits in sympathetic modulation of cardiac and arterial functions with aging occur in the presence of exaggerated neurotransmitter levels. Plasma levels of norepinephrine and epinephrine, during any disturbance from the supine basal state, increase to a greater extent in older compared with younger healthy humans. The age-associated increase in plasma levels of norepinephrine results from an increased spillover into the circulation and, to a lesser extent, reduced plasma clearance. The degree of norepinephrine spillover into the circulation differs among body organs; increased spillover occurs within the heart. Deficient norepi nephrine reuptake at nerve endings is a primary mechanism for increased spillover during acute graded exercise. During prolonged exercise, however, diminished neurotransmitter reuptake might also be associated with depletion and reduced release and spillover. Cardiac muscarinic receptor density and function are also diminished with increasing age and might contribute to the decrease in baroreflex activity observed in aged subjects. 43

CARDIOVASCULAR RESERVE IN THE PRESENCE AND ABSENCE OF A Ş-ADRENERGIC BLOCKER



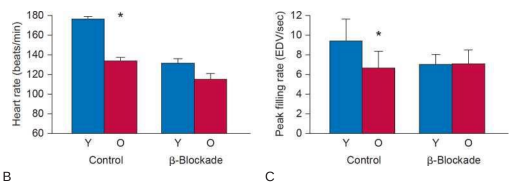


FIGURE 25-3 Cardiovascular reserve in the presence and absence of a beta-adrenergic blocker.

25

Structural Change	Functional Change	Mechanisms	Molecular Mechanisms
Myocyte size	Prolonged contraction	Prolonged cytosolic Ca2+ transient	
Myocyte number	Prolonged action potential	i SR Ca⊶ pumping rate i Pump site density i I _{Ca} inactivation	i SR Ca⊶ pump mRNA No change in calsequestrin mRNA T Na-/Ca- exchanger mRNA
	Diminished contraction velocity	i Iro density i a-MHC protein T p-MHC protein i Myosin ATPase activity i RXRpI and Y mRNA i RXRpI and Y protein i Thyroid receptor protein	i a-MHC mRNA T B-MHC mRNA No change in actin mRNA i RXR01 and Y mRNA
	Diminished beta-adrenergic contractile response	i Coupling beta-adrenergic receptor-acyclase No change in Gractivation No change in BARK activity i TNI phospholamban i Phospholamban phosphorylation i Ica augmentation i Ca transient augmentation T Enkephalin peptides T Proenkephalin mRNA	i Beta-adrenergic receptor No mRNA change in BARK mRNA
Matrix connective tissue	T Myocardial stiffness i Growth response i Heat shock response	T Hydroxyline proline content T Activity of myocardial RAS T Atrial natriuretic peptide	T Collagen mRNA T Fibronectin mRNA T ATIR mRNA T Atrial natriuretic peptide mRNA T Induction of immediate-early genes i Activation of HSF

AT₁R, angiotensin AT₁ receptor; BARK, beta-adrenergic receptor kinase; Ca, intracellular calcium concentrations; G, inhibitory G protein; HSF, heat shock factor; I_a, calcium influx; MHC, myosin heavy chain; mRNA, messenger RNA; RAS, renin-angiotensin system; RXR, retinoid X receptor; SR, sarcoplasmic reticulum; TNI, troponin I.

Cardiac Aging in Animal Models

Cardiac Structure

Cellular and molecular mechanisms involved in age-associated changes in myocardial structure and function have been largely studied in rodents. Some underlying mechanisms that drive agerelated cardiac remodeling are listed in Table 25-3. The altered a significant driver of increased oxidative stress. Notably, NADPH cardiac structural phenotype that evolves with aging in rodents includes an increase in LV mass due to an enlargement of myocyte size and focal proliferation of the matrix in which the myocytes reside, which may be linked to an altered cardiac fibroblast number forces. 51 or function. The number of cardiac myocytes becomes reduced because of necrosis and apoptosis, with necrosis predominating.

neighboring myocytes. 44 Stretch of cardiac myocytes and fibroblasts initiates growth factor signaling (eg, angiotensin II/TGF-B), which, in addition to modulating cell growth and activation of the calcineurin-NFAT pathway increases with age. 45 is increased in the senescent rodent heart.

There is evidence that transcriptional events associated with hypertrophic stressors become altered with advancing age; for example, the nuclear binding activity of the transcription factor nuclear factor- KB is increased and that of another transcription factor, Sp1, is diminished. A proteomic comparison study 47 oxygen species (ROS) in many different tissues. 49

Angiotensin II Signaling

Many experimental studies have shown that increased activation of the renin-angiotensin-aldosterone system (RAAS) is a prominent feature of age-related cardiac remodeling that may account for many of the phenotypic changes observed. 8.50

It is increasingly appreciated that RAAS activation may itself be oxidase family members are activated by angiotensin II, aldosterone, endothelin 1, cytokines (eg, TNF- a), growth factors (eg, TGF- \$, platelet-derived growth factor), and mechanical

The regulated production of small amounts of ROS in response to such stimuli appears to be ideally designed for involvement in Putative stimuli for cardiac cell enlargement with aging in redox signaling pathways, and an involvement of NADPH rodents include an age-associated increase in vascular load due to oxidases in such signaling has indeed been demonstrated in many arterial stiffening and stretching of cells caused by dropout of studies. 51 Experimental studies have confirmed that NADPH oxidase-derived ROS play important roles in the pathogenesis of cardiovascular pathological processes such as endothelial dysfunction, atherosclerosis, diabetic vasculopathy, RAAS-related matrix production, leads to apoptosis. In the mouse heart, hypertension, and ischemic vascular remodeling. Recently, it has also been shown that NADPH oxidase activation is involved in The expression of atrial natriuretic 46 and opioid peptides, several types of cardiac remodeling, including angiotensin IImolecules that are usually produced in response to chronic stress, induced cardiac hypertrophy, aldosterone-induced cardiac fibrosis, and cardiac remodeling after myocardial infarction. 52-54 In addition, increased myocardial NADPH oxidase activity is found in human heart failure. 55,56

In male Fischer 344 cross brown Norway rats aged 2 months (young rats), 8 months (young adult rats), or 30 months (old rats), age-associated increases in blood pressure, cardiomyocyte area, discovered 117 proteins to be differentially expressed in the aged coronary artery remodeling, and cardiac fibrosis were associated left ventricle. An increase in oxidative stress is also implicated in with increased myocardial NADPH oxidase activity, which was age-related cardiac remodeling, 48 and aging is indeed well attributable to the Nox2 isoform. These changes were accompanied recognized to be associated with increased production of reactive by increased expression of connective tissue growth factor and TGF- 1 and by a significant activation of matrix metalloproteinase 2 (MMP-2) and MT1-MMP, a membrane-bound activator of MMP-

were chronically treated with angiotensin II for 28 days. Other function. cytokines that could be involved in the process include TNF -a, which is implicated in aging-related disease possibly by an and pharmacological scenarios that increase Ca 2 influx (eg, upregulation of MMP levels. 57 TNF -a is also a known activator of neurotransmitters, postischemic reperfusion, or oxidative stress). 62 NADPH oxidase, and its possible involvement in aging-related In hearts or myocytes from the older heart, enhanced Ca 2 influx, cardiac remodeling merits investigation.

including angiotensin II, aldosterone, endothelin 1, and cyto kines, aforementioned age-associated adaptation =" that occurs within the it is arguable that they may be a suitable therapeutic target for the cells of senescent heart (and also of young animals chronically prevention of aging-related cardiac remodeling in addition to exposed to arterial pressure over load). The amounts and ratios of targeting of the individual agonists (such as the RAAS) that activate each class of proteins involved in cell ion homeostasis, the lipid the oxidase.

Calcium Regulation

Age-altered myocardial Ca 2 - cycling is closely related to coordinated changes in levels of sarcoplasmic reticulum (SR) Ca 2+ -handling proteins or their function. In aged hearts, selective downregulation of sarcoendoplasmic reticulum calcium-ATPase Increased Mitochondrial ROS Generation fl in the Aged Heart During Acute Stress phosphorylation (which might be reflected in the desensitized adenylyl cyclase response), and increased levels of Na ·/ Ca ² · Mitochondria possess a unique Ca ² · transport machinery, regulate old ventricle similar to those of the young organism. 58

Cardiac myocyte Ca ²-cycling is modulated by beta-adrenergic receptor stimulation. The well-documented age-associated changes that may enhance their sensitivity and promise their receptor to adenylyl cyclase through the G s protein and changes in myocardium are modified or impaired by aging. adenylyl cyclase protein. This leads to a reduction in the ability to sufficiently augment cell cyclic adenosine mono phosphate and to mechanism, and this protection is reduced in the older heart. This activate protein kinase A to drive the phos phorylation of key may relate to decreased heat shock protein HSP-70 expression. proteins that are required to augment cardiac contractility. In Nitric oxide, produced by endothelial nitric oxide synthase (eNOS) contrast, the apparent desensitization of beta-adrenergic receptor or inducible nitric oxide synthase (iNOS), plays a role in signaling with aging does not appear to be mediated by increased cardioprotection. However, iNOS and eNOS expression and beta-adrenergic receptor kinase or increased G, activity.

peptide receptor signaling because of the significant antagonistic specific sites targeted by each protein kinase C isoform. effects between stimulation of opioid peptide receptor and betaadrenergic receptor-mediated positive contractile response. 59

Remarkably, the available examinations of cardiac excitationcontraction coupling in rodents of both sexes 60 or females show stiffening caused by interlocking of actin and myosin). 62 Although apparent increase in the SR Ca²-content, (2) the lack of the effect of mitochondria contribute the the increase in the SR Ca ² content on the configuration of the Ca ² · transients and contraction, and (3) a reduction of fractional SR Ca ²·release, in the presence of unchanged I _{CaL}. Nevertheless, these results provide an initial direct demonstration of sexually dimorphic changes 423 in cardiac excitation-contraction coupling associated with normal adult aging.

Calcium Regulation During Acute Stress. Aging-associated major changes in heart structure and function place the aged heart constantly "on the edge" and at risk, including the loss of its adaptive response to stress. Acute excess myocardial Ca 2-loading leads to dysregulation of Ca 2-homeostasis, impaired diastolic and systolic function, arrhythmias, and cell death. 61 The cell Ca 2+ load is determined by membrane structure and permeability characteristics, the intensity of stimuli that modulate Ca 2-influx or

efflux by their impact on the regulatory function of proteins within The changes in old rats were replicated in 8-month-old rats that membranes, and ROS, which affect both membrane structure and

Excessive cytosolic Ca 2 · loading occurs during physiological impaired relaxation, and increased diastolic tone occur during Because NADPH oxidases are activated by multiple factors, pacing at an increased frequency. 63 This is a "downside" of the milieu of mem branes in which these proteins reside (eg, types and amounts -j-- of long-chain polyunsaturated acids), and the threshold fattiness required to produce ROS and ROS-derived alkenes all change with aging and reduce the threshold for acute Ca²·overload that occurs within the older heart.

exchanger could lead to the reduction in SR Ca 2 · load. It is also metabolism under partial control by Ca 2 · , 64 and may themselves suggested that the aged heart uses the compensatory increase in the contribute to cell Ca 2-homeostasis. Mitochondrial production of L-type Ca 2 - currents and the significant prolongation of action ROS in the heart and mitochondrial dysfunction associated with potential duration to preserve SR loading and to keep the reperfusion appear to increase with age. 65,66 Thus, mitochondria amplitude of intracellular Ca 2 transients and contractions in the appear to play a central role in the reduced Ca 2 tolerance observed in aged animals.

reduction in the postsynaptic response of myocardial cells to beta-response to stress (such as ischemia) but can also result in the loss adrenergic receptor stimulation appears to be the result of multiple of the ability to trigger cardioprotective mechanisms. These changes in the molecular and biochemical steps that couple the changes can obviously have deleterious effects on the fate of the receptor to postreceptor effectors. The major limiting modification mitochondrion, cell, and organism. Compared with young adult of acute beta-adrenergic receptor signaling with advancing age in myocardium, the senescent myocardium is more sensitive to rodents appears to be at the coupling of the beta-adrenergic ischemia, 67 suggesting that protective pathways existing in adult

Repetitive ischemia is an endogenous cardioprotective activity are modified in senescent myocardium. 68 Changes in The age-associated reduction in beta-adrenergic receptor protein kinase C translocation have been described in aged signaling may, in part, also be related to upregulation of opioid myocardium, 69 leading to certain age-associated changes in the

Several studies indicate that reperfusion that follows ischemia generates ROS and that exogenous application of ROS to cells causes Ca 2 · overload, ATP depletion, and rigor (ie, cellular only a lack of age-related changes in the configuration of the Ca 2. multiple enzymes including NADPH oxidase at the plasma transient in myocytes from female hearts. The somewhat membrane and cyclooxygenases and xanthine oxidase in the unexpected findings in aged female myocytes include (1) an cytoplasm also contribute to the overall oxidative burden,

oxidative phosphorylation.

conductance channel in the mitochondrial inner membrane that catastrophe." causes the loss of the proton motive force in the respiratory chain 25 the MPT induction can lead to high [CaJ and to the generation of inactivation of a -ketoglutarate dehydrogenase during reperfusion additional ROS, whereby necrosis and apoptosis are initiated and has been identified. 73 cell death ensues. Studies have discovered a phenomenon referred to as ROS-induced ROS release, ⁷⁰ that is, the initial or "trigger" ROS peroxidation. Indeed, omega-6 rather than omega-3 PUFAs appear generated by mitochondria in response to laser excitation of to be preferred targets of ROS-induced peroxidation that produces fluorescent mitochondrial membrane label leads to an HNE. Thus, the increase in omega-6/omega-3 PUFA that occurs amplification of ROS production by the mitochondria themselves. with aging may be a mechanism for increased HNE production

collapse of the mitochondrial membrane potential, which renders attenuated by an omega-3 PUFA-rich diet. Notably, an omega-3 the mitochondrial membrane permeable to bulk movement of ions, PUFA-rich diet also prevents the age-linked decrement of the larger molecules, and water between the cytosol and the mitochondria-specific membrane phospholipid cardiolipin, a mitochondrial matrix. Cells from senescent hearts have a crucial cofactor for cytochrome- c oxidase and adenine nucleotide substantially lower threshold for the generation of ROS-induced translocase (ANT) activity. 74 ROS release and less likelihood of the MPT. ROS, such as the at or near the site of their formation.

mouse hearts is manifested by disrupted cristae and vacuolation enhanced. 75 (loss of electron density) and is accompanied by an increase in mtDNA copy number and by increased mitochondrial transcription factor A, nuclear respiratory factors, and PPAR y change in LV mass or myocardial dysfunction. Furthermore, mitochondrial biogenesis. 45

Reduction in Membrane Polyunsaturated Fatty Acids

Membrane polyunsaturated fatty acids (PUFAs) undergo lipid older heart, the response in many instances is reduced. peroxidation by ROS, producing various aldehydes, alkenals, and nonenal (HNE). 71 HNE, potentially the most reactive of these contribute to the pattern of gene expression observed in the hearts compounds, is formed by superoxide reactions with membrane of senescent rodents and may also dictate the limits of adaptive omega-6 PUFAs and reacts with protein sulfhydryl groups to responses to the imposition of additional chronic stress. The acute induce altered protein conformation. 71 Because the mitochondrial induction of both immediate-early genes and later responding respiratory chain represents a major subcellular source of ROS genes that are expressed during the hypertrophic response is during reperfusion of ischemic myocardium, and because blunted in hearts of aged rats after aortic constriction. 77 Similarly, ischemia-reperfusion has been associated with decreased rates of the acute induction of HSP-70 genes in response to either ischemia NADH-linked, membranes are likely targets of lipid peroxidation and HNE- of adaptive capacity is observed in younger rats that have used a mediated enzymatic dysfunction. The concentration of HNE is part of their reserve capacity before a growth factor challenge. 79 increased in the reperfused postischemic myocardium. 72

can therefore diffuse from the site of its origin in membranes to models of pressure overload and myocardial infarction. 41 Sexually affect potential targets distant from the initial site of ROS dimorphic cardiac phenotypes have also been production. 71 Thus, if membrane bilayer PUFAs are converted to lipid hydroperoxides, then lipid peroxidation may be viewed as an "amplifier" for the initial ROS. Furthermore, the reactive aldehydes generated in this process may well act as "toxic second messengers" of the complex chain reactions that follow ROS production. 71 Ischemic HNE modification of mitochondrial proteins occurs exclusively in hearts isolated from senescent rats, but it has not

majority of ROS generation as a byproduct of electron transfer and been established whether ROS-induced peroxidation of mitochondrial membranes leading to the production of HNE is The permeability transition pore (PTP) is a nonselective, high-related to the amplification of ROS and "mitochondrial

Studies show that exposure of intact cardiac mitochondria to and the failure to generate ATP. ROS generated within HNE leads to decreased NADH-linked respiration, partly due to mitochondria play a role in causing the occurrence of HNE-dependent inactivation of a -ketoglutarate dehydrogenase mitochondrial membrane permeability transition (MPT). The and pyruvate dehydrogenase. These enzymes contain lipoic acid release of mitochondrial contents into the cytosol resulting from residues that are prime targets for HNE. An age-associated

PUFAs differ with respect to their susceptibility to lipid This amplification is associated with induction of the MPT and after ischemia-reperfusion, but this effect can be markedly

ANT may participate in the formation of a nonspecific hydroxyl radical (OH), are highly reactive and generally short- membrane pore through a Ca 2 · -mediated, cyclophilin Dlived species. Therefore, they might be expected to cause damage dependent conformational change in ANT. 75 Cyclophilin D binding to ANT increases after oxidative stress that enhances the Mitochondrial ROS lead to mitochondrial DNA damage and Ca 2 - dependent formation of the PTP, which induces MPT. dysfunction in a feed-forward manner, causing functional cellular Adenine nucleotides located in the mitochondrial matrix bind to and organ declines. Protein carbonyls and oxidative damage are ANT and decrease the sensitivity of the PTP to Ca²·, but this effect increased in mitochondrial extracts of the aged mouse heart, and is antagonized by modification of specific thiol groups on ANT – mitochondrial DNA point mutation and deletion frequencies by oxidative stress products such as HNE or by thiol reagents such increase approximately threefold. 45 Mitochondrial damage in as carboxyatractyloside – to the extent that MPT induction may be

Reduced Chronic Adaptive Capacity of the Older Heart

Many of the multiple changes in cardiac structure, excitation, coactivator 1 a (PGC-1 a). 45 Life span is prolonged by more than myofilament activation, contraction mechanisms, Ca 2 · dys-15% in mice with catalase overexpression targeted to the regulation, deficient beta-adrenergic receptor signaling, and mitochondria, and these mice do not exhibit an age-associated altered gene expression of proteins involved in excitation contraction coupling that occur with aging (see Table 25-2) also catalase overexpression results in significant reductions in occur in the hypertrophied myocardium of younger animals with mitochondrial oxidative damage, mtDNA mutation deletion experimentally induced chronic hypertension 76 and in failing frequencies, mitochondrial protein carbonyls, and activation of animal or human hearts, in which they have been constructed as an adaptive response to a chronic increase in LV loading. When chronic mechanical stresses that evoke substantial myocardial hypertrophy (eg, pressure or volume overload) are imposed on the

Transcription factors that influence expression of a number of hydroxyalkenals, such as malonaldehyde and 4-hydroxy-2- genes become altered in abundance during aging and may ADP-dependent respiration, mitochondrial or heat shock is reduced in hearts of senescent rats. 78 A similar loss

Premature development of heart failure or death in males In contrast to many ROS species, HNE is rather long-lived and compared with matching females has been documented in rat

general, transgenic models of heart failure present a more rapid onset or a greater severity of cardiac dysfunction in male versus female hearts. For instance, mice with cardiac specific Despite the fact that age is the dominant risk factor for overexpression of TNF- exhibited heart failure and increased hypertension and atherosclerotic disease, most prior clinical

without dilation and only a small reduction of basal LV fractional associated vascular changes were not identified and in part due to shortening and response to isoproterenol; male mice showed a the view that whatever changes were present could not be large LV dilation, reduced fractional shortening relative to both prevented or treated. During the past 10 years, however, there has wild-type littermates and transgenic females, and minimal been considerable progress in understanding the vascular changes response to isoproterenol. 81 Cardiac myocyte hypertrophy was associated with age and how they may be modified. Some of these similar in male and female transgenic mice. Compared with wild- interventions are currently used only for those with demonstrated type mice, myocytes from female TNF- a transgenic mice displayed disease or "abnormal" blood pressure or lipid levels. Preclinical and a slower decline of the Ca 2 transient but similar amplitudes of Ca some small clinical trials suggest, however, that they may be isoprotere nol. In contrast, the amplitude and the rate of decline of vascular properties before the development of hypertension or Ca² transients and contractions and the response to isopro terenol clinically evident vascular disease. This section discusses those were significantly reduced in myocytes from male transgenic TNF- currently available interventions that may modify these processes. a mice. 81

Myocyte Progenitors in the Aging Heart

Observations in humans and animals suggest that myocyte maturation and aging are characterized by loss of replicative potential, telomeric shortening, and expression of the senescenceassociated protein/cell cycle inhibitor p16 INK4a. 82 84 Telomeric shortening in Physical conditioning status is strongly associated with precursor cells leads to the generation of progeny that rapidly acquire the cardiovascular events and mortality. 91 It is probable that one senescent phenotype. The activity of telomerase, an enzyme responsible mechanism is an impact on the age-associated vascular present only during cell replication, was decreased 31% in aging changes that contribute to the marked increase in hypertension and male rat myocytes but increased 72% in female counterparts. 85 atherosclerosis with advanced age. Thus, understanding of the biology of cardiac precursor cells, including factors enhancing the activation of the precursor cell vascular stiffness. 92 Studies in middle-aged and the relatively pool, their mobilization, and translocation, may facilitate the "younger" older groups indicate that aerobic exercise can favorably development of novel strategies to prevent or to reverse the affect central vascular stiffness. Tanaka and associates 93 employed diminished adaptive capacity to increases in pressure and volume both cross-sectional and intervention studies to assess the impact loads (and perhaps heart failure) in the old population. This of physical conditioning status, and changes in status, on central involves progressive increase in the size of the cell (up to a critical arterial compliance using carotid B-mode ultrasonography and volume beyond which myocyte hypertrophy is no longer possible); applanation tonometry in men without disease as assessed by deficits in the electrical, Ca 2-cycling, and mechanical properties; history, examination, laboratory studies, electrocardiography, and, and cell death.

Cardiac myocytes with senescent and nonsenescent phe pressure and volume loads. 87,88

by an imbalance between factors enhancing oxidative stress, intervention was performed in 20 previously sedentary men during telomere attrition, and death and factors promoting growth, 3 months. It consisted primarily of walking, an average of 5 migration, and survival. Recent findings suggest a preeminent days/week for 42 minutes/day, at 73% of maximum heart rate. position of IGF-1 among factors that interfere with cardiac cellular Although maximal oxygen consumption did not change with this senescence. Specifically, cardiac-restricted overexpression of IGF-1 regimen, there was a 25% increase in central arterial compliance, in transgenic mice has been shown to delay the aging myopathy to a level not significantly different from that in the enduranceand the manifestations of heart failure 89 and to restore SERCA-2a trained men in the cross-sectional study. Thus, a relatively short expression and rescue age-associated impairment of cardiac and modest exercise program can significantly affect the important myocyte contractile function. 90 This effect was also partially age-associated increase in central vascular stiffness. mimicked by short-term in vitro treatment with recombinant IGF-1. 90 Furthermore, intramyocardial delivery of IGF-1 improved direct measures of stiffness are reported in both men and women. senescent heart phenotype in male Fischer 344 rats, 83 including - 94,95 In addition to affecting systolic and pulse pressures , vascular increased proliferation of functionally competent precursor cells compliance also interacts with ventricular stiffness and contractility and diminished angiotensin II-induced apoptosis. Myocardial to affect cardiac output. One proposed index of this ventricularregeneration mediated by precursor cell activation attenuated arterial coupling, or interaction, is the beat-to-beat relationship ventricular dilation and the decrease in the ratio of ventricular mass between LV end-diastolic to chamber volume, resulting in improvement of in vivo cardiac function in animals at 28 to 425 29 months of age. 83

discovered in some studies in genetically engineered mice. 41,40 In INTERVENTIONS TO RETARD CARDIOVASCULAR **AGING**

mortality that is markedly higher in young males than in females studies in older individuals focused on evaluating and (~50% and 4%, respectively, by 20 weeks of age). 80 implementing interventions that targeted "traditional" risk 25 At 12 weeks of age, female mice displayed LV hypertrophy factors (eg, hypertension and elevated lipids), in part because age-² · transients and contractions and the inotropic response to beneficial in retarding or ameliorating age-associated altered

Exercise and Vascular Risk Factors

One of the most important of these is an increase in central in those older than 40 years, exercise testing.

In the cross-sectional study, central arterial compliance was notypes already coexist at young age. 86 However, aging limits the approximately 40% higher in the endurance-trained (participation growth and differentiation potential of precursor cells, thus in vigorous aerobic exercise > 5 times/week) than in the sedentary interfering not only with their ability to sustain physiological cell or recreationally active (light to moderate exercise > 3 times/week) turnover but also with their capacity to adapt to increases in older groups. The best correlate of arterial compliance in all groups was maximal oxygen consumption (r = 0.44 to 0.45), probably the The loss of precursor cell function with aging is mediated partly best index of physical conditioning status. The exercise

Similar observational, cross-sectional study results using less

changes in intrathoracic pressure. 96

relationship, the index was significantly higher, indicating 110 Physical activity is also associated with improved mood in showed a trend to improved coupling with the exercise have not been examined. intervention. 97 The benefits of aerobic exercise are less apparent 25 in octogenarians. 98 In one study of 22 men and women (mean months of exercise training are associated with significant age, 82 ± 4 years), 9 months of training at higher than 80% of peak improvement in vascular structure and function, at least in the heart rate did not change arterial stiffness as assessed by carotid "younger" group of older individuals (ie, in the seventh decade). applanation tonometry.

distension-induced modification of collagen cross-linking and progenitor cell number in healthy older individuals 112 or in terms vascular smooth muscle relaxation due to enhanced nitric oxide of carotid intima media thickness. In addition, the majority of the signaling or decreased sympathetic tone. In addition to the exercise studies were performed in men, and it is unclear whether favorable influences on systolic blood pressure associated with similar benefits occur in women. Importantly, the clinical relevance decreased stiffness (less of a change in pressure for any given of the findings is unclear; although stiffness and the other studied ejected stroke volume), chronic physical activity also likely vascular parameters are associated with important cardiovascular enhances baroreceptor-mediated parasympathetic cardiac stimuli, outcomes, it is unclear whether exercise-induced changes in these 99 which may maintain vascular homeostasis during hypervolemic variables actually change these outcomes. Finally, the effects of and hypovolemic stresses as well as decrease the likelihood of different types, duration, and intensity of exercise on vascular adverse cardiac arrhythmias.

The type of exercise training may, however, influence the effect beneficial effects. on central vascular stiffness. 98 In contrast to aerobic exercise, and although it has not been examined in older adults, one intervention on the basis of comorbid conditions and baseline fitness levels, study in young men 100 and another in young women 101 which may limit the type and duration of exercise . Because of this demonstrated that high-intensity resistance training significantly variability, a subjective intensity of exercise is recommended by the increased arterial stiffness, an effect that was reversed after a American College of Sports Medicine and the American Heart deconditioning period. 100 A third intervention study using a more Association concerning the initiation, type, and intensity of conventional resistance protocol, again in a young population, physical activity in older Americans. 113 Thus, moderate-intensity showed no effect. 102

endothelial function, ^{103,104} believed to be related to several factors, activity is recommended for 20 minutes a day, 3 days each week. most importantly decreased nitric oxide bioavailability, Moderate is defined as 5 or 6 and vigorous as 7 or 8 on a 10-point accompanied by decreased prostacyclin and increased endothelin scale; sitting is 0. Moderate activity produces "noticeable increases" clinical consequences are believed to include increased produces large increases in heart rate and breathing. Resistance or

studies in previously sedentary middle-aged and older men be 5 to 8, where 0 is no movement and 10 is maximum effort. indicate that aerobic exercise training can prevent and reverse, Additional goals include exercises directed to maintenance and respectively, age-associated declines in endothelial function. 105,106 strengthening of flexibility and balance. Several mechanisms are likely to be involved. The initiating event is most likely increased shear stress resulting from the increased pulse pressure during exercise. 107 This leads to an acute increase in nitric oxide but also to chronic changes that increase nitric oxide Premenopausal bioavailability, including increased eNOS gene expression, atherosclerotic disease than do men, premature menopause is reduced oxidative stress, and decreased production and increased associated with increased vascular risk, and observational studies scavenging of free radicals. 84,108

exercise training may result in favorable arterial remodeling , postmenopausal women 114 and in numerous pre-clinical models resulting in decreased shear stress for any given exercise-induced and studies 115 suggest significant benefits. These include increase in stroke volume. The importance of shear stress is stimulation of eNOS in vascular endothelial cells, 116 improved indicated by studies reporting that exercise of large muscle groups mitochondrial function and decreased oxidative stress in vascular results in improved endothelial function in distant arteries (eg, endothelium, 117 reduced angiotensin II-induced free radical improved forearm endothelial function after leg exercise), whereas production in vascular smooth muscle cells, and stimulation of exercise of small muscle groups (eg, one forearm) does not have endothelial progenitor cell production 118 and the catalytic subunit remote effects. The extent of endothelial function improvement of human telomerase. 71 All with exercise training is directly related to the extent of preexisting dysfunction, suggesting that even healthy older individuals would benefit. In general, benefits are evident as early as 4 to 8 weeks after the initiation of training and are lost within several weeks after cessation of an exercise program.

There are additional potential although less investigated favorable effects of exercise on vascular risk in older individuals. In one study, 109 a 3-month aerobic exercise program decreased endothelin 1-mediated vasoconstriction in 15 older men, mean age

pressure and stroke volume as modified by respiratory-induced 62 ± 2 years; and in another, endothelial-mediated tissue-type plasminogen activator in human forearms was increased by 55% in In a study examining the impact of physical fitness on this 10 men, mean age 60 ± 2 years, after 3 months of aerobic exercise. improved coupling, in Master athletes (mean age, 68 ± 3 years) adults with depression, a risk factor for vascular disease. 111 The than it was in older sedentary individuals; in an intervention relationships among changes in physical activity, increased mood, study, a 1-year period of endurance training in nine individuals decreased anxiety, and vascular function and structure, however,

In summary, habitual exercise and periods as short as weeks to Despite these favorable effects, these changes are not uniform; for Potentially responsible mechanisms include mechanical example, there appears to be no benefit in terms of endothelial function require additional study, as do the mechanisms for the

Specific activity recommendations are tailored to the individual aerobic physical activity is recommended for a minimum of 30 Another important age-related vascular change is a decrease in minutes a day, 5 days each week; or vigorous-intensity aerobic 1, systemic angiotensin II, and sympathetic vasoconstrictors. The in heart rate and breathing, and vigorous-intensity activity development and progression of atherosclerosis and inappropriate muscle strengthening activities should be performed on two more vasoconstriction, which may result in acute coronary syndromes. nonconsecutive days per week with a weight that allows 10 to 15 Cross-sectional observational studies as well as intervention - repetitions for each major muscle group. The level of effort should

Estrogen and Vascular Risk Factors

women experience indicate that estrogen use is associated with decreased vascular -Sustained increases in nitric oxide associated with long term risk. The effects of estrogen on coronary endothelial function in

of these correlate well with endothelial cell senescence and aging, and the favorable impact of estrogen on them would suggest a significant clinical benefit.

Nevertheless, the Women's Health Initiative, the most recent and largely regarded as definitive study of estrogen in postmenopausal women with prior hysterectomy, demonstrated no benefit on the primary composite endpoint of non- fatal HMG-CoA Reductase Inhibitors myocardial infarction or death and an increase in stroke and pulmonary embolism. 119 Although it was concluded that estrogen should not be used for primary or secondary prevention in postmenopausal women, as is currently recommended, studies of the potential benefits of this therapeutic approach still continue. Therapy was started in the Women's Health Initiative study an average of 10 years after menopause, when significant changes in the vascular substrate had already occurred.

The preclinical studies all suggest a preventive effect of estrogen, rather than a reversal of preexisting abnormalities, arguing that estrogen administration during the perimenopausal or early postmenopausal period should be studied. Furthermore, polymorphisms of the estrogen receptor are known to significantly alter the vascular response to estrogen administration 120 and cardiovascular disease risk, suggesting that genome-based estrogen therapy may be beneficial. In addition, new selective estrogen receptor modulators may provide greater vascular benefit and less vascular risk than current estrogen preparations.

Testosterone and Vascular Risk Factors

There is considerable epidemiological evidence that androgen deficiency in older men is associated with increased cardio vascular mortality and atherosclerotic risk factors. 121,122 The most important function, and survival have been extensively studied. Statins of these risk factors are the components of the metabolic syndrome, increase endothelial progenitor cell numbers by increasing bone including increased waist-to-hip ratio, insulin resistance, and marrow hematopoietic stem cells and inducing endothelial hypertension, but they also include increased total and low-density lipoprotein cholesterol and carotid intima-media thickness and decreased flow-mediated dilation.

associated with hypertension, insulin resistance, and vascular - the impact of age as a risk factor for vascular disease and the stiffening. Small studies in hypogonadal older men indicate that benefits of statins on the responsible mechanisms, randomized androgen treatment can increase both flow-mediated and trials examining the impact of statins on age changes in vascular nitroglycerin-mediated brachial artery vasodilation , decrease substrate and relating them to subclinical and clinical systolic and diastolic blood pressures, improve lipid profiles, and manifestations of atherosclerosis appear warranted. reduce inflammatory cytokines. The extent to which these may be mediated, in part, by their favorable effect on body composition, Renin-Angiotensin-Aldosterone Inhibition including increased lean body and muscle mass and decreased visceral and total body adiposity, is not known.

Despite these small studies as well as strong basic and preclinical plausibility for a beneficial effect of androgen based therapies in patients with increased cardiovascular risk and low testosterone levels, there are relatively few placebo-controlled younger and middle aged groups. randomized trials. The conduct of such trials is limited by several factors. Testosterone may be associated with increased risk for goals of less than 140 mm Hg versus 130 mm Hg in more than 1100 prostate hypertrophy and acceleration of subclinical prostate

Oral therapies may be associated with liver toxicity and injectable forms with discomfort and inconvenience. In this regard, selective androgen receptor modulators may be a desirable alternative. These may be taken orally without liver toxicity and in animal models demonstrated positive effects of androgen therapy while eliminating some negative effects. As any treatment would be most effective in those with hypogonadism, the determination of whether an individual patient is hypogonadal is not mean testosterone levels, there is considerable individual variability. In this regard, administration of androgen therapy to those with normal levels may

result in no benefit and unwanted side effects, whereas 427 administration of placebo to those with low testosterone is not ethical. Free and bioavailable rather than total testosterone levels may most clearly correlate with any androgen- mediated benefit and risk.

The beneficial effects of statin therapy on cardiovascular events in older subsets of randomized trials are similar to those in the younger groups. 123,124 In the Heart Protection Study subset of more than 5800 individuals 70 years or older who were randomized to placebo or simvastatin, the occurrence of nonfatal myocardial infarction, coronary death, stroke, or revascularization was significantly lower in the group randomized to statin (23.6%) than in the group randomized to placebo (28.7%) during the 5-year study period.

The pleiotropic effects of statins, however, are remarkably suited to inhibit the adverse consequences of aging on the vascular substrate that predisposes to the development, progression, and clinical manifestations of atherosclerosis. These include increased nitric oxide bioavailability, anti inflammatory effects, improved endothelium-dependent vasoreactivity, reversal of vascular endothelial growth factor-induced endothelial hyperpermeability, 125 and probable stabilization of vulnerable atherosclerotic plagues. One of the most important aspects of the vascular substrate that is altered with age concerns endothelial progenitor cells, which affect vascular function, integrity, and, most importantly, repair.

The effects of statins on endothelial progenitor cell number, progenitor cell differentiation through the phosphatidylinositol 3kinase/Akt pathway, improve progenitor cell functional capacity, 126 and enhance endothelial progenitor cell survival and prevent Androgen deprivation therapy in men with prostate cancer is telomere erosion in the presence of oxidative stress. 127 Considering

The importance of hypertension as a risk factor for vascular disease and mortality 128 and the benefits of antihypertensive therapy 129,130 in the older population are well documented. In fact, the benefit of intervention is at least as great in the older population as it is in the

A randomized study compared treatment systolic pressure nondiabetic individuals with a systolic pressure of 150 mm Hg or higher. 131 In the subset of patients older than 70 years, the odds ratio of experiencing the primary outcome (development of left ventricular hypertrophy, a potent predictor of cardiovascular events) in the lower versus the higher systolic pressure goal group was 0.49, whereas that for the entire study cohort was 0.61.

In addition to arteriolar changes, which increase peripheral resistance, changes in the central vasculature associated with aging (eg, fibrosis, inflammation, and media thickening) may predispose to atherosclerosis and vascular events, 7 some of which may be straightforward. Despite the significant age-associated decrease in particularly related to increased angiotensin II activity. As such, these changes offer additional targets that, if successfully treated, may decrease event rates over and above those achieved with blood pressure lowering alone.

428 Although nearly all antihypertensive therapies can lower blood pressure in older individuals, those that inhibit the RAAS may also ameliorate these age-specific changes. Thus, ACE inhibition decreases measures of central vascular stiffness in human studies, in part independent of blood pressure reduction, ^{25,132} as well as TNF -a , a mediator of inflammation. ¹³³ Angiotensin receptor blockade also has favorable effects on indices of central vascular stiffness, including pulse wave velocity and aortic augmentation index, as well as the media-to-lumen ratio measured from 25 biopsy specimens. ¹³⁴

Although ACE inhibitors and angiotensin receptor blockers have different sites of action, and the reduction in arterial stiffness is significantly greater when the combination is used, ¹³⁵ a large randomized study (ONTARGET) demonstrated adverse clinical outcomes with the combination compared with the individual agents. ¹³⁶ In this trial, there was a 9% increase in the primary outcome (dialysis, doubling of creatinine, or death) and a 24% increase in dialysis or creatinine doubling in the patients randomized to ramipril and telmisartan compared with those randomized to ramipril alone.

Aldosterone antagonists are useful in the management of blood pressure in individuals with primary hyperaldosterone ism who will not undergo surgery; in many patients with resistant hypertension, defined as inability to achieve blood pressure goal despite the use of three antihypertensives; and in those with hypertension and hypokalemia. ¹³⁷ Older individuals may be more likely to experience hyperkalemia and therefore require closer monitoring, and sex hormone-related side effects can be largely avoided with the use of eplerenone, a more selective agent.

Apart from the diuretic effect, aldosterone antagonists have additional benefits that may improve age-associated vascular changes. Aldosterone increases vascular inflammation , perivascular fibrosis, 138 and vascular expression of the angiotensin type 1 receptor. In addition, individuals stiffness in with hyperaldosteronism is greater than what would be expected on the basis of blood pressure alone. 139 Treatment with aldosterone antagonists prevents the ageassociated increase in arterial stiffness and fibrosis in old, normotensive rats by altering vascular collagen and elastin. In patients, aldosterone antagonism decreases pulse wave velocity and augmentation index measures of central vascular stiffness, 140 and in a separate study, it decreased stiffness in gluteal subcutaneous resistance vessels as well as several circulating inflammatory mediators. 141 Thus, inhibition of the RAAS not only lowers blood pressure and improves outcomes in older individuals with hypertension but also reduces those ageassociated vascular changes that predispose to the development of hypertension and atherosclerosis. The risks associated with relying on surrogate markers alone, however, are well illustrated by the results of the ONTARGET study, which demonstrates again the importance of obtaining meaningful clinical outcomes before any general recommendations concerning antiaging strategies can be advised and implemented.

Tuxedo

Although the proportion of older individuals who smoke is lower than that of the rest of the population, 10.8% of men and 8.4% of women older than 65 years do so. ¹⁴² There is consistent evidence that cigarette use is associated with an increased risk of adverse cardiovascular outcomes in all age groups. In a prospective data analysis of more than 1 million adults,

the relative risk for ischemic heart disease, cardiovascular disease, and mortality was increased above that of those who never smoked by approximately 1.5-fold in the subset of those who were older than 65 years. 143 The potential mechanisms are multifactorial, including increased ROS, altered autonomic tone, platelet activation, increased arterial stiffness, and probably many of the toxins in the product itself. 144 These would affect several of the changes associated with the mechanisms underlying increased atherosclerotic risk associated with vascular aging. Exposure to cigarette smoke impairs endothelial function as assessed by flow-dependent brachial artery vasodilation, increases elaboration of vascular endothelial growth factor (one response to vascular injury), decreases nitric oxide production, and decreases endothelial progenitor cell function as assessed by chemotaxis. 145 Some of these vascular effects are reversed with cigarette cessation, 146 and cessation is associated with decreased risk for cardiovascular events as well. 147 Cigarette cessation should be encouraged for all populations of patients.

Considerations in Older Individuals

The increased prevalence of atherosclerotic risk factors, disease, and complications once an event has occurred emphasizes the importance of risk factor identification and aggressive, appropriate intervention in the older population. Primary and secondary prevention strategies do not differ between the older and younger populations at risk for or with cardiovascular disease. As noted before, similar or more significant clinical benefits (because of the increased absolute risk) accrue with lipid- and blood pressure-lowering therapy , at least up to 80 years of age. Cigarette cessation and other lifestyle factors, particularly maintaining physical activity, are valuable from cardiovascular and multiple other health perspectives.

However, age changes in body composition and in renal and hepatic function and the increased likelihood of comor bidities and concomitant medications may influence the type of therapy selected, the intensity of the regimen, the dose of the agent, and the dosing interval. Age is associated with increased bleeding risk with antiplatelet and anticoagulation therapies. Low doses, attempts to avoid combining anti platelet agents, and careful monitoring are all more important in the older age groups. Advanced age is often associated with a decrease in lean muscle and an increase in fat body mass. Renal function as defined by formulas for creatinine clearance includes age as one variable.

Age is associated with a decrease in hepatic clearance of drugs as well, particularly those metabolized through the cytochrome P-450 pathway. Older individuals may also be more likely to experience side effects, such as constipation, urinary retention, postural hypotension, and cognitive changes, because of preexisting impairments or concomitant medications. Excellent resources are available, including those from the Food and Drug Administration, regarding drug selection, dosing, and interactions in all age groups.

CLINICAL IMPLICATIONS CARDIOVASCULAR AGING

OF

Our present understanding of the age-associated alterations in cardiac and arterial structure and function at both the cellular and molecular levels provides valuable clues that may assist in the development of effective therapies to prevent, to delay, or to attenuate the cardiovascular changes that accompany aging. Many of these age-associated changes are being increasingly recognized as risk factors for cardiovascular diseases. In spite of the interest in the physiology of the age-associated changes in cardiovascular structure and function, cardiovascular aging has

remained, for the most part, outside of mainstream clinical medicine. This is because the pathophysiological implications of these changes are largely under appreciated and are not well disseminated in the medical community. In fact, age has traditionally been considered a

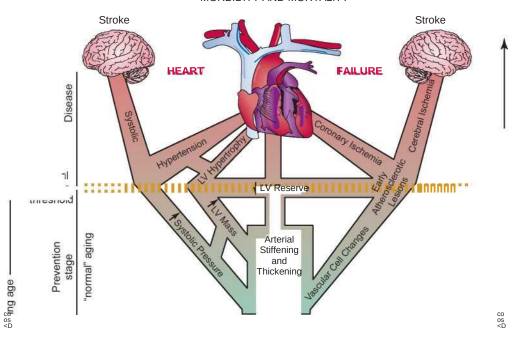


FIGURE 25-4 Aging: the major risk factor for cardiovascular morbidity and mortality.

Clinical practice threshold

non-modifiable risk factor. Because many of the age-associated alterations in cardiovascular structure and function, at both the cellular and molecular levels, are specific risk factors for cardiovascular diseases (see Table 25-3), there is an urgency to incorporate cardiovascular aging into clinical medicine.

One way to conceptualize why the clinical manifestations and the prognosis of these diseases worsen with age is that in older individuals, the specific pathophysiological mechanisms that cause clinical disorders are superimposed on heart and vascular substrates that are modified by aging (Fig. 25-4). Imagine that age increases as one moves from the lower to the upper part of Figure 25-4 and that the line bisecting the top and bottom parts represents the clinical practice "threshold" for disease recognition. Thus, entities above the line are currently classified as "diseases" and lead to heart and brain failure.

The vascular and cardiac changes currently thought to occur as a result of the "normal aging process" (ie, those addressed in the previous sections) are depicted below the line. These age-associated changes in cardiac and vascular properties alter the substrate on which cardiovascular disease is superimposed in several ways. First, they lower the extent of disease severity required to cross the threshold that results in clinically significant signs and symptoms. For example, a mild degree of ischemia-induced relaxation abnormalities that may be asymptomatic in a younger individual may cause dyspnea in an older individual, who, by virtue of age alone, has preexisting slowed and delayed early diastolic relaxation.

Åge-associated changes may also alter the manifestations and presentation of common cardiac diseases. This usually occurs in patients with acute infarction, in whom the diagnosis is delayed because of atypical symptoms resulting in increased time to onset of therapy. Age-associated changes, including those in beta-adrenergic responsiveness and in vascular stiffness, also influence the response to and therefore the selection of different therapeutic interventions in older individuals with cardiovascular disease.

In one sense, those processes below the line in Figure 25-4

ought not to be considered to reflect normal aging. Rather, they might be construed as specific risk factors for the diseases that they relate to and thus might be targets of interventions designed to decrease the occurrence or manifestations of cardiovascular disease at later ages. Such a strategy would thus advocate the treatment of normal aging. Additional studies of the specific risks of each normal age-associated change are required.

CONCLUSION

Aging is the dominant risk factor for cardiovascular diseases. However, aging should no longer be viewed as an immutable risk factor. A steady stream of incremental knowledge, derived from both animal and human studies, has established that several of the aging-associated changes in the heart and in the walls of the central arteries are themselves potent and independent risk factors for cardiovascular diseases. This suggests that these age-associated alterations in arterial and cardiac structure and function could represent the link that explains, at least in part, the risky component of aging.

Policy makers, researchers, and clinicians should intensify their efforts towards identification of novel pathways that could be targeted for interventions aiming at retardation or attenuation of these age-associated alterations, particularly in individuals in whom these alterations are accelerated. Future studies would then examine whether these strategies (ie, those targeting cardiovascular aging) can have a salutary impact on the adverse cardiovascular effects of accelerated cardiovascular aging. As such, cardiovascular aging is a promising frontier in preventive cardiology that is ripe for and in dire need of attention.

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25



Diagnostic Testing to Help Improve Risk Prediction

CHAPTER 26

Concepts of Screening for Cardiovascular Risk Factors and Disease

KEY POINTS
Donald M. Lloyd-Jones

- Screening involves the routine testing of asymptomatic individuals for the purpose of detecting the presence of a condition or a disease. The ultimate goal of screening is to identify the disease or condition of interest in an early phase, when intervention may be more effective in reducing subsequent morbidity or mortality.
- A number of metrics are available to assess the performance of screening tests, including sensitivity, specificity, predictive values, model fit characteristics, discrimination measures, and risk reclassification.
- Unlike with cancer, screening for CVD has typically involved screening for risk factors, for which there is solid evidence of utility, rather than for CVD itself.
- Current CVD clinical practice guidelines recommend screening for global cardiovascular risk by use of multivariable risk equations to assist in decision making regarding the intensity of prevention strategies.

 Use of imaging modalities to screen for subclinical CVD is an area of intense research interest. Current guidelines do not recommend routine screening for atherosclerosis in asymptomatic individuals, but accumulating

Screening involves the routine evaluation or testing of asymptomatic individuals for the purpose of detecting the presence of a condition or a disease. The ultimate goal of screening is generally to identify the disease or condition of interest in an early, or latent, phase, when intervention may be more effective in reducing subsequent morbidity or mortality. Historically, many of the concepts that have been used to define the utility of screening tests (sensitivity, specificity, predictive values) in patients and populations arose from the assessment of tests designed to detect medical conditions with a clear pathological diagnosis, such as cancer. In recent years, these classical measures of diagnostic utility and many novel metrics have been employed as screening increasingly is used for preclinical conditions in the causal pathways for clinical disease and for prognosis, rather than merely for diagnosis. In the realm of cardio vascular disease (CVD), screening is a topic of significant import and heated debate, given the high incidence of disease across the life span, the substantial burden of

evidence may help define a role for these screening tests, perhaps in refining risk stratification among those at intermediate risk for CVD events by traditional risk factor levels

morbidity and mortality, and the potentially high costs of screening tests and therapies. This is especially true in the current health care and economic environment. Thus, understanding of the basic concepts related to screening for CVD is of paramount -importance. A review of the conceptual frame work for screening tests and the means for evaluating their utility will serve as -background for a discussion of the relative merits of screening for CVD risk factors (-traditional and novel), global cardiovascular risk, and subclinical atherosclerosis.

CONCEPTS IN SCREENING

Screening Criteria

In 1968, the World Health Organization ¹ defined criteria for screening for diseases in medicine. As shown in Box 26-1, these criteria indicate that screening for disease should be considered when the disease has a significant impact (in terms of prevalence or severity), has an adverse natural history

3OX 26-1 World Health Organization Criteria fo Screening

The condition sought should be an important health problem for the individual and community.

There should be an accepted treatment or useful intervention for patients with the disease.

The natural history of the disease should be adequately understood.

There should be a latent or early symptomatic stage.

There should be a suitable and acceptable screening test or examination.

Facilities for diagnosis and treatment should be available.

There should be an agreed policy on whom to treat as patients. Treatment started at an early stage should be of more benefit than treatment started later.

The cost should be economically balanced in relation to possible expenditure on medical care as a whole.

Case finding should be a continuing process and not a once and for all project.

Modified with permission from Wilson JMG, Jungner G: Principles and practice of screening for disease. WHO Chronicle 22:473, 1968.

that is well understood, and is treatable or modifiable during its asymptomatic phase. Of equal importance in these criteria are the features of the screening test itself: it should be reliable , available, cost-effective relative to other strategies, and applicable in an ongoing fashion. These widely accepted criteria provide a useful framework for evaluation of new (and old) screening tests. Clearly, it is inadequate merely to use a test with face validity for detection of disease; rather, careful consideration of the potential benefits, risks, harms, and potential costs associated with screening must be undertaken before widespread clinical adoption.

Types of Screening

Mass or universal screening involves assessment of all individuals in a population or group (eg, all school-aged children or all pregnant women). Case finding, or high-risk screening, involves the application of a screening test to sub groups identified as being at higher risk than average because of the presence of known risk factors (eg, a strong family history of disease). As discussed later, each of these approaches has merit, depending on the nature of the disease or condition being screened and the knowledge of important predisposing factors.

Assessment of Screening Tests

A list of commonly applied metrics for evaluation of screening tests in CVD is provided in Table 26-1. Some discussion of these tests is warranted for a fuller understanding of their implications. Classical metrics of test characteristics (sensitivity, specificity, and related metrics) can be understood most easily in comparison with a "gold standard" test indicated in the definitive presence or absence of disease. However, they are also applicable to prognostic tests, in which the gold standard is the development (incidence) of disease during the follow-up interval of observation after testing.

As recently detailed by the American Heart Association, ² appropriate consideration of traditional and novel screening tests for CVD (which may include single tests or multivariable risk scores) should entail assessment of a number of different metrics beyond simple association, sensitivity, specificity, and predictive values. Demonstration that a screening test has a significant statistical association with the outcome of interest is necessary but clearly not sufficient for evaluation of its utility. A number of metrics are available to assist in the evaluation of the

performance and utility of risk estimation models. These metrics assess characteristics of the test (similar to a diagnostic test), its ability to discriminate cases from non-cases, the calibration of the model, model fit, and the informativeness of the model for the outcome of interest. Newer methods of assessment, such as analysis of risk reclassification , also allow comparison of different risk stratification algorithms by use of novel markers or risk scores. Knowledge of a few of these metrics and concepts will suffice for most clinicians to interpret the utility of risk prediction models. Consideration of all of these factors is important to understand the utility of a risk score.

Sensitivity, Specificity, and Predictive Values

As shown in Figure 26-1 and Table 26-1, sensitivity and specificity reflect the true-positive and true-negative rates, respectively. In other words, a test with high sensitivity will detect a large proportion of individuals who have disease; a test with high specificity will correctly be negative in individuals without disease. These are useful test characteristics that in most cases do not change on the basis of the prevalence of disease in the groups being tested. However, they do not necessarily answer the question that is of interest to a clinician and patient: Is disease present? Positive and negative predictive values typically may be more useful as assessments of diagnostic and screening tests because they indicate the likelihood of having or developing disease given a positive (or higher) or negative (lower) test result, but their heavy reliance on the incidence and prevalence of disease in the population may make them difficult to translate from one clinical scenario to another.

Measures of Model Fit and Informativeness

Other measures, such as the Bayes information criterion, are now commonly used to assess the utility of statistical risk prediction models that include screening tests. These tests can indicate whether a risk model is predicting disease incidence better than chance alone. They can further indicate whether the addition of new screening tests to a base model provides better risk prediction than the base model alone, provided all of the same individuals are being assessed by both models.

Discrimination

One of the most widely reported measures of model discrimination for screening tests in general, and CVD risk prediction models specifically, is the area under the receiver operating characteristic curve (AUC), or C-statistic. The Cstatistic is a function of both the true-positive and false-positive rates of the screening tool across all of its values, and it represents the ability of the score to discriminate (future) cases from noncases. In other words, the C-statistic indicates the probability that a randomly selected patient who has or develops the disease (a "case") will have a higher test result or risk score than a randomly selected non-case. The AUC or C-statistic can vary from 1.0 (perfect discrimination) to 0.5 (random chance, equivalent to flipping a coin to determine case status). Thus, a C-statistic of 0.75 for a given model would indicate that a randomly selected case has a higher score than a randomly selected non-case 75% of the time (Fig. 26-2). C-statistics below 0.70 are generally considered to indicate inadequate discrimination by a test, whereas those between 0.70 and 0.80 are considered "acceptable," and between 0.80 and 0.90, "excellent." 3

The C-statistic is imperfect as a stand-alone metric for assessment of screening tools or risk prediction models. In general, the C-statistic indicates whether a test or risk score is generating appropriate rank-ordering of risk for cases and



I ABLE 26—1 Commonly Applied	Measures and Terms for Assessment of the Utility of Screening Tes	sts *
Measure	Definition	Comment
Sensitivity	Proportion of those with disease who have a positive test result $[P(T + D +)]$	Detection rate, true-positive rate; high sensitivity is useful for ruling out disease
Specificity	Proportion of those without disease who have a negative test result $[P(T \longrightarrow D \longrightarrow)]$	True-negative rates; high specificity is useful for ruling in disease
False-negative rates	Proportion of those with disease who have a negative test result [P(T — D +)]	Equal to 1 — sensitivity; when low, useful for ruling out disease
False positive rate	Proportion of those without disease who have a positive test result $[P(T + D -)]$	Equal to 1 — specificity; when low, useful for ruling in disease
Positive predictive value (PV +)	Proportion of those with a positive test result who have disease $[P(D + T +)]$	Dependent on the prevalence (incidence) of disease
Negative predictive value (PV —)	Proportion of those with a negative test result who do not have disease [P(D — T —)]	Dependent on the prevalence (incidence) of disease
Likelihood ratio	Ratio of true-positive rate to false-positive rate $[P(T + D +)] / [P(T + D -)]$	Can be used to calculate post-test odds (and therefore post-test probability of disease) by multiplying with pretest odds
Pretest probability	Probability of disease based on available information (prevalence, or risk-adjusted prevalence)	
Post-test probability	Adjusted probability of disease after application of additional (screening) test	
Area under the receiver operating characteristic curve (AUC, or C-statistic)	Function of true-positive rate and false-positive rate across all values of diagnostic (screening) test	Indicates the discrimination ability of the test; likelihood that a randomly selected case will have a positive (or more adverse) test result compared with a randomly selected non-case
Fit model	Assessment of whether statistical model including test improves case detection/prediction compared with chance or base model	Information criteria or likelihood ratio test often used to assess the utility of the model
Calibration	Degree to which screening (prediction) test or model accurately predicts absolute levels of observed event rates	Usually assessed with the Hosmer-Lemeshow test
Net reclassification improvement		
	Degree to which new test increases predicted risk (across a decision threshold) for those who subsequently have events and decreases predicted risk (across a decision threshold) for those who do not subsequently have events	
Integrated discrimination index	It indicates how far individuals are moving, on average, along the continuum of predicted risk after application of the test	Equivalent to the difference in R - between the two models being compared

[·]In the case of screening tests related to prognosis (rather than to diagnosis), the definitions would be relevant to those who do or do not develop disease during observation. P, probability; D+, disease present; D—, disease absent; T+, test positive; T—, negative test.

	Disease present	Absent disease	
Positive test	A (True positive; TP)	b (False positive; FP)	a + b
Negative test	c (False negative; FN)	d (True negative; TN)	c + d
	a + c	b + d	a + b + c + d

Sensitivity = True-positive rate = $P(T+ \mid D+) = TP/(TP+FN) = a \ I \ (a+c)$ Specificity = True-negative rate = $P(T- \mid D-) = TN/(TN+FP) = b \ I \ (b+d)$ Positive predictive value = $P(D+ \mid T+) = TP/(TP+FP) = a \ I \ (a+b)$ Negative predictive value = $P(D- \mid T-) = TN/(TN+FN) = d \ I \ (c+d)$

FIGURE 26-1 Calculation of utility measures for diagnostic or screening tests.

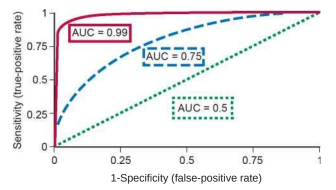


FIGURE 26-2 Representative curves depicting the area under the receiver operating characteristic curve (AUC or C-statistic).

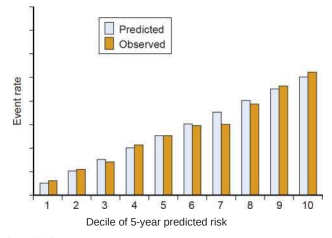


FIGURE 26-3 Assessment of calibration of a risk score or prediction test by comparing predicted 5-year risk with observed event rates, stratified by decile of predicted risk.

non-cases, not whether the predicted risk and observed outcome rates are similar (which is a function of calibration) or how much greater the estimated risk for disease is between selected cases and non-cases. $^{4.5}$

Pepe and coworkers ⁶ have demonstrated that very large odds ratios (or relative risks) are required to reach meaningful levels in the C-statistic. For example, a univariate odds ratio of 9.0 or greater would be required to achieve a C-statistic that provides excellent discrimination of cases from non cases for a continuous screening test (eg, cholesterol or coronary calcium) in which the distribution of test scores differs by 2 or more standard deviations. These magnitudes of differences in distribution of test scores are rarely seen in clinical practice (where risk factor levels often overlap substantially), as are such high odds ratios. However, the combination of multiple, independent screening tests or risk markers, as in the Framingham risk score (FRS) and similar scores (see later), does provide these magnitudes of relative risk.

Calibration

Measures of calibration assess the ability of a screening test or risk prediction model to accurately predict the absolute level of risk that is subsequently observed. Demonstration that a risk prediction model is well calibrated would require that if the model estimates that the risk for a certain subgroup of individuals is 5% during 5 years, then the observed event rate should be close to 5%. Calibration is often assessed visually by dividing the population at risk into strata, such as deciles of predicted risk, and plotting the predicted risk versus the observed event rate for each decile (Fig. 26-3). The statistical metric used to test for the calibration of a risk model most often is the Hosmer-Lemeshow x 2 test. A *P* value < 0.05 for such a test would indicate poor calibration of the model for the population.

Assessment of Appropriate Risk Reclassification

A newer paradigm to assess the utility of screening tests and risk prediction models, which is recommended by the American - Heart Association ² for appropriate assessment of such tests, is risk reclassification analysis. ⁷ This approach requires measurement of the proportion of individuals who are reclassified from one risk stratum (eg, intermediate risk) based on the estimated risk provided from a first model to a different risk stratum (eg, high risk) based on estimated risk from a model that contains the additional test information. Some of these risk reclassifications end up being appropriate (based on subsequent observed events): some individuals who have events are reclassified to higher predicted risk strata, and some who do not have events are reclassified to lower predicted risk strata. However, some reclassifications are inappropriate, moving

future cases to lower predicted risk strata and future non-cases to higher predicted-risk strata.

Pencina and coworkers ⁷ have proposed two indices, the net reclassification improvement (NRI) and the integrative discrimination index (IDI), to attempt to quantify the appropriateness and the amount of overall reclassification. In general, the NRI indicates how much more appropriate reclassification occurs than inappropriate reclassification with use of the new model. The NRI can vary from - 2, indicating that all individuals are reclassified inappropriately, to + 2, indicating that all are reclassified appropriately. In other words, if the newer test reclassifies all of the people who have events upward, and all of the people who do not end up with events downward, the NRI would be + 2. For this test, a P value < 0.05 suggests that significantly greater number are being reclassified appropriately than are being reclassified inappropriately. The IDI can be thought of as indicating how far individuals are reclassified, on average, along the continuum of predicted risk. 8 If the IDI is small (even if it is statistically significant), then a given individual's change in predicted risk with the new model will be small, on average. As an example, consider a new risk prediction model or test that is being compared with the FRS for stratifying a population into risk categories. The new model might have a significant NRI, reclassifying a net of 10% of people more appropriately; but if the IDI is small (eg, less than 1%), then most of the net reclassification is occurring immediately adjacent to the decision thresholds that separate the risk categories, such as a change from a predicted risk of 19.8% with an old model to a predicted risk of 20.4% with a new model. This change might cross the decision threshold for treatment, but it would indicate no real impact in understanding or forecasting the patient's risk, especially if the decision thresholds are relatively arbitrary. The significance of such small movements is also heavily dependent on the threshold selected. Indeed, this scenario is what is often observed in current studies comparing older and newer CVD risk prediction scores. 9

Interpretation of Risk Information Provided by Screening Tests

Different types of information about risk for disease may be garnered from screening tests. The relative risk of disease is the ratio of disease or disease incidence among those with a positive test result compared with those who have a negative test result. As such, relative risk measures the strength of the association between the test and disease, but relative risks are poor indicators of clinical utility, and physicians and patients often have difficulty interpreting relative risk estimates 10-12 in the absence of an obvious comparison group. A relative risk for disease of 10 might seem very high, but if the incidence rate in the referent group is close to 0, it will also be close to 0 in the group with the relative risk of 10. Absolute risk of disease is often expressed as the estimated rate of development of new cases of disease per unit of time (or incidence) in individuals with a positive test result. Absolute risk is more easily understood than relative risks, and they allow clinical recommendations for interventions in individuals who exceed unacceptable risk thresholds. 10-12 This approach has been widely adopted for 5and 10-year estimation of absolute risks for coronary heart disease (CHD) and CVD to guide clinical decision making for CVD prevention. ¹³ The attributable risk of a test result describes the proportion of the incidence of disease in a population associated with the test result, assuming a causal relationship exists. The population attributable risk takes into account the proportion

of individuals in the population who test positive as well as the relative risk. Therefore, attributable risk is a useful concept in selecting screening tests that might be targeted for prevention

In clinical trials, we often consider the concept of a number needed to treat (NNT) to prevent one event. The NNT is calculated as follows:

1 (rate of disease in the control group - rate of disease in /c the **intervention** group

Thus, an intervention that reduces mortality from 7% to 5% during 5 years would have an NNT of 1 / (0.07 - 0.05) = 1 / (0.02)= 50. In other words, 50 people would need to be treated with the intervention for 5 years to prevent one event.

A similar concept, the number needed to screen (NNS), has been proposed. 14 Equivalently, the NNS represents the number of people who would have to be screened to detect one person with disease (as in mammography for breast cancer detection or faecal occult blood testing for colon cancer) or one person with a level of a risk marker that is deemed to be unacceptably high (eg, detection of one individual with an Agatston coronary artery calcium score > 300). The NNS is calculated as follows:

1/(rate of disease in the usual care group - rate of disease in / (the scree **n** ed group or

1 (prevalence of risk marker level greater than threshold in)/(a

given gr o up

The NNS may be best calculated from clinical trial data that compares a screening strategy with usual care, but it can be derived from observational data as well. The concept has been expanded to adjust for willingness (or lack thereof) to undergo screening in certain groups, thus indicating a number needed to be invited for screening, which is generally a higher number among lower risk healthier populations, and a number actually needed to screen, a lower number among potentially motivated, higher risk screenees. 15,16

Law ¹⁷ has pointed out several weaknesses of the NNS. First, there may be difficulty generalizing the NNS from one population to another that may have a different basal rate of disease. Similarly, it may be impossible to compare the NNS for two different screening modalities or two different diseases if they are performed in different populations. NNS estimates are also sensitive, meaning that small changes in factors that influence disease rates may have a large impact on NNS. In addition, events are not only prevented; some are postponed, which may also have significant value, but it is difficult to capture in shorter term studies. 17 Nevertheless, the NNS may represent a useful construct if appropriate comparisons are made.

Biases and Pitfalls of Screening Tests

Whereas it may seem intuitive that screening for disease would always be useful, a number of pitfalls related to screening are routinely encountered in clinical practice. It is thus important to determine objectively whether screening can ultimately improve outcomes.

Imperfection of Screening Tests: False-Positives, False-Negatives, and Implications

The discussion of metrics for assessment of screening tests introduced the concepts of false-positive and false-negative results of screening tests. False-negative results may be dev astating, delaying diagnosis of a potentially deadly disease or condition. If screening tests are selected for maximum sensitivity , this eventuality may be minimized. The implications of falsepositive results are more subtle but potentially just as important because they will affect much larger numbers of individuals. If a

disease is uncommon, with a prevalence of < 50% in the 437 population to be screened, it is virtually certain that there will be more false-positive than true-positive test results. For example, a seminal paper in the literature of screening mammography examined the 10-year risk of false positive mammograms. 18 Based on data from 2400 women aged 40 to 69 years at entry, a group for whom mammography has consistently been shown to reduce the risk of breast cancer deaths, there were an average of four mammograms per formed during 10 years. Of the women who were screened, the false-negative rate was approximately 1%, but 24% had at least one false-positive mammogram, 13% had at least one false-positive breast examination, and a total of 32% had at least one false-positive result for either test, requiring further evaluation. Thus, a large proportion of women without breast cancer underwent further imaging or biopsy with associated anxiety, pain, potential harms of unnecessary procedures, and substantially increased costs due to falsepositive results. The cumulative risk of a false-positive result after 10 mam mograms was estimated to be 49%. 18 The issue of false-positives is further complicated when screening tests are used for prognosis because the gold standard is the occurrence of an event, which may not have happened (yet) within the observed follow-up but could happen soon thereafter, changing a false-positive into a true-positive.

Selection Bias, or Spectrum Bias

As discussed earlier, the utility of a screening test is dependent on the population being tested. For example, exercise treadmill testing is known to have higher sensitivity (ie, higher detection rate) among people with three-vessel or left main coronary disease than among people with one-vessel coronary disease. 19 Thus, screening with treadmill testing will appear better among those selected to be at very high risk (or who may have unrecognized symptoms) than if it is applied to a more general healthy population. In addition, although one may invite entire groups or populations for screening, some individuals will be more likely than others to be screened. If people with a higher risk of disease are more likely to be screened, then the screening test could appear worse than it really is because adverse outcomes among the screened population will be higher than for the general population. 15 Conversely, if a test is selectively available to younger, healthier individuals, the screened group will have lower disease rates than the general population and will appear to have benefited from screening.

Adherence/Compliance Bias

Compared with the general population, people who are more likely to adhere to prescribed therapies may also be more willing to undergo screening, therefore improving the apparent impact of the screening strategy in preventing adverse outcomes.

Lead Time Bias

The effectiveness of a screening test may be overestimated because it detects disease at a much earlier stage than would otherwise have been the case. Therefore, the time between "diagnosis" of disease and death will be prolonged, on average, compared with diagnosis at the time of symptom onset. However, if the outcome of early detection is similar to the outcome with symptomatic detection, this lead-time bias will give a false sense of effectiveness to the screening test: "survival with disease" is artifactually lengthened by earlier disease detection, even though age at death may be the same for both groups. In the meantime, the patient may



438 have experienced anxiety or economic hardship related to the diagnosis. The amount of lead-time bias is a function of the sensitivity of the screening test as well as of the biological rate of disease progression.

Length-Time Bias

Length-time bias has been best conceptualized in the frame work of cancer detection, but it may have significant harm to subclinical CVD detection as well. Many screening tests may perform better in detecting slower growing cancers and that have a better prognosis than in detecting rapidly growing cancers because of a longer preclinical or latent phase in the slow-growing cancer. If true, then screening will tend to detect more cancers that would not have killed the patient or 26 may not even have been symptomatic before death from other causes. In parallel, it is possible that some CVD screening tests could detect stable Given all of these potential pitfalls and biases in evaluating screening clinically manifested.

Prevalence Bias

A related concept is prevalence bias, in which early rounds of screening will tend to detect slower growing cancers, with good prognosis, whereas subsequent rounds of screening will detect newer, more rapidly growing cancers. This bias should be considered in the context of defining screening intervals.

Overdiagnosis Bias

Screening may also detect benign or harmless conditions that interest. mimic the disease of interest. This may make the screening strategy appear to be more effective than it actually is because the harmless condition would not have resulted in an adverse outcome. However, it would likely have led to unnecessary RISK FACTORS treatment.

Combining Screening Tests

Use of multiple screening tests leads to several issues. First, they may be correlated; significant amounts of correlation and test will be more useful.

Diagnostic Test Cutoffs

One final issue that deserves consideration is the selection of for diagnostic or screening tests. The optimal cutoff point is a RISK function of sensitivity and specificity, the prevalence of stream testing.

Presentation of Risks and **Benefits of Screening**

A substantial literature has addressed the issues related to the presentation of data regarding the risks and benefits of screen ing. These issues include problems of framing risk estimates, Framingham Assessments with presentation of relative risk reductions, which may seem large, versus absolute risk reductions, which are usually very screening may sound large, particularly for a scary disease like 0.75 to 0.80. 30-34 The FRS and similar risk scores breast cancer, but this represents only 1 or 2 deaths prevented or postponed for 1000 women screened annually for 10 years. The "average" amount of life gained for a woman in her 40s is thus 3 days, ²⁰ but it might be decades for the few women who are identified with breast cancer. Other problems include physicians' and patients' perceptions of risk, their ability to assess the benefits and harms related to screening, and what

value they might place on those outcomes, among many

However, if there exists a safe, inexpensive, and reasonable therapy that can provide substantial risk reduction, these problems may be surmountable by lowering the threshold for treatment substantially with little potential for harm. In such an environment, the implications of a false-positive result may be minimized. It is easy to suggest that we would rather overtreat than undertreat in such a situation. This paradigm may now be driving CVD screening, with the increasing availability of potent, generic statin medications.

Appropriate Evaluation and Application of **Screening Tests**

subclinical CVD better than they detect CVD that will become tests and strategies, it is imperative that new screening modalities be assessed rigorously before widespread application. This is particularly true for costly screening modalities. The best data come from randomized controlled clinical trials of screening strategies in appropriate populations across the spectrum of health and risk, with blinded evaluation of outcomes and appropriate gold standard or outcome definition. Subsequent analyzes can then examine the relative and absolute reduction in disease, the number needed to screen (or harm), and the effectiveness and cost effectiveness of the screening strategy in the general population or specific subgroups of

CARDIOVASCULAR SCREENING

Formal assessment of the risks, benefits, harms, and costs of screening for dyslipidemia, hypertension, and diabetes have been performed. 21-26 Although some data are lacking, acknowledgment that these are causal risk factors for CVD, the lack of independence may lead to false certainty when results available evidence on the ability of these risk factors to identify are similar. For most screening situations, sequential testing individuals at heightened risk for CHD, and the substantial data will be preferred to use of concurrent multiple screening tests indicating efficacy, effectiveness, and cost effectiveness of therapy because the results from a first test may be sufficient, or they (certainly for dyslipidemia and hypertension) have led to strong may revise the pretest probability to a point where a second recommendations for universal screening among middle-aged and older adults and generally accepted practice of screening younger adults. ²¹⁻²⁶ These are unlikely to be revised in the future.

appropriate thresholds (cutoffs) to define a positive test result SCREENING FOR GLOBAL CARDIOVASCULAR

disease in the population to which it will be applied, and any Current clinical practice guidelines from the US National benefits of correct diagnoses as well as risks and costs of Cholesterol Education Program (ATP III), 13 international guidelines, incorrect diagnoses with regard to potential harm and down ^{27,28} and American Heart Association guidelines ²⁹ have adopted and thoroughly incorporated the strategy of "global" risk estimation using multivariable equations containing traditional risk factors as a means to identify those who should receive more intensive therapy. The US Preventive Services Task Force (USPSTF) has supported the use of global risk estimation as a screening strategy for the selection of patients warranting lipid-lowering therapy. 23

Risk Score

small for screening tests. Thus, a relative risk reduction of 15% In most cohorts studied to date, the FRS provides very good in breast cancer mortality associated with mammographic - discrimination, as indicated by C-statistics that typically range from

discriminate risk better for women than they do for men, often ethnic groups. 30 having C-statistics in excess of 0.80 for women. In addition, the FRS has good discrimination in most populations (including those from prediction, a number of investigators have examined the addition of outside the United States) in which it has been studied because it multiple markers simultaneously to traditional risk equations. In the contains age as a covariate and because of the fairly universal Cardiovascular Health Study, I Shlipak and associates 40 reported associations between CVD and the major traditional risk factors the that addition of six novel biomarkers (including interleukin-6, CRP, FRS also incorporates.

Americans, and native Chinese, discrimination remains acceptable, but the FRS tends to overestimate risk. 35,36 However, with simple assess reclassification as well. In a cohort of 5067 Swedish steps to recalibrate the FRS model, it performs well in both individuals free of CVD at baseline, observed for a median of discrimination and calibration. 35

provided by existing multivariable risk prediction models, they two independently significant novel biomarkers were added to remain the logical standard to which new risk markers or subclinical the traditional models, the C-statistics improved by 0.007 and disease measures from screening tests must be added to demonstrate 0.009, respectively, for CVD and CHD events, indicating improvement or against which newer models must be compared. minimal clinical utility. Likewise, the proportion of participants The methods that should be used to evaluate novel markers of reclassified was modest (8% overall for CVD risk and 5% for cardiovascular risk were recently summarized and recommended CHD risk). The NRI was 0 for CVD events and 0.047% for CHD by a special panel of the American Heart Association. 2 All of the events, indicating no or only minimal net improvement in risk metrics discussed before should be examined and assessed for reclassification. Greater improvements in reclassification were statistically significant improvement, and then clinical judgment observed in analyzes restricted to intermediate-risk individuals, applied to determine whether there is also clinically meaningful - but any correct reclassification was almost entirely due to down improvement that would affect decision making in a reasonable ward reclassification of individuals who did not experience number of patients. That said, if the change in the C-statistic is small events (yet). for a new one compared with an existing risk score, or when a novel risk marker is added to an existing risk score, it is extremely unlikely for CVD (including CRP, leukocyte count, fasting blood that the novel risk score or marker will provide a clinically glucose, lipoprotein(a), and homocysteine) and made no meaningful improvement as a screening tool across the population. recommendation for use of these markers because of However, the novel risk marker may still add value within certain insufficient evidence for potential benefits and harms. 42 The subgroups. To demonstrate these concepts, it will be useful to American Heart Association and the Centers for Disease consider some published examples.

Attempts to Improve Risk Prediction Through Addition of Novel Markers to Existing Risk Scores

A substantial body of literature during the last 5 years has been devoted to examining the addition of newer risk markers to traditional risk scores. Essentially all of the single additional markers There now exists a large number of published risk prediction studied to date have yielded little additional clinical benefit when scores for various CVD or CHD endpoints that have been considered as screening tests across the entire spectrum of risk. For derived from a variety of different populations. One recent example, Folsom and colleagues 37 observed minimal changes in the example, the Reynolds risk score for women, 9 included the C-statistic (0.000 to 0.011) for prediction of CHD events using 19 traditional risk factors plus CRP levels and parental history as different novel bio markers added individually to the FRS. However, selected for inclusion in the model based on improvement in the within subgroups predicted to be at intermediate risk by traditional Bayes information criterion. Compared with a Framingham-like models, the addition of new risk markers can help reclassify some model recalibrated for this study's endpoint, overall individuals, and this is often the group for which the addition of discrimination and calibration of the two models were similar. information from a new test is most clinically useful. In the Women's Internal validation of the Reynolds risk score revealed that 5.8% Health Study, 31 when C-reactive protein (CRP) was added to a of 8149 women were reclassified when the Reynolds risk score model with traditional risk factors, the C-statistic did not improve was applied (compared with the FRS), with about half of those measurably for the whole population (the reported C statistic was being reclassified upward and half down ward. The clinical 0.81 for both models). However, women in the middle of the implications of downward reclassification are unclear because predicted risk spectrum in this cohort (FRS predicted risk of 5% to withholding of therapy from those potentially at higher risk on 9%) with a CRP level > 3.0 had an observed event rate that was equal the basis of traditional risk factors does not seem warranted to or greater than that of some women with a FRS-predicted risk of given available clinical trial data. The > 10%. However, the traditional risk model (FRS) determines a far greater magnitude of risk than does the level of CRP, indicating the 440 generalizability and utility of this score and many others important context provided by the traditional risk model. 38 Similar results have been observed with

addition of parental history information to traditional risk 439 scores, 39 with minimal change in overall discrimination but potential benefit in intermediate-risk groups. More recent data suggest a larger change in the C-statistic associated with the addition of coronary artery calcium scores to the FRS, with increments in the C-statistic from 0.02 to 0.11 in different racial and

Because single risk markers have not added substantially to risk fibrinogen, lipoprotein(a), and factor VIII levels plus presence of Calibration of the FRS in various populations may differ anemia) | to a traditional risk model improved the C-statistic from substantially, however, because of various CVD incidence rates in 0.73 26 to 0.74 in older individuals with chronic kidney disease but different settings. Studies to date indicate that the FRS discriminates decreased the C-statistic from 0.73 to 0.72 in those with normal renal CHD risk very well and is well calibrated for a wide range of white function. In the Framingham study, the addition of brain natriuretic and African American populations in the United States. For other peptide levels plus microalbuminuria to a traditional risk model for populations, such as Asian Americans, American Indians, Hispanic CVD events yielded an increase in the C-statistic from 0.76 to 0.77. 41

More recent studies have extended these types of analyzes to 12.8 years, models with traditional risk factors had C-statistics Given the overall acceptable to excellent risk discrimination - of 0.758 and 0.760 for CVD and CHD events, respectively. When

> The USPSTF recently evaluated nontraditional risk markers Control and Prevention have recommended against routine screening with CRP or other inflammatory markers but concluded that it might be reasonable to aid in decision making for patients at intermediate predicted risk for CHD. ⁴³

Examples of Newer Risk Prediction Models

are still being assessed.

Limitations of 10-Year Risk Prediction Models

Ten-year risk estimation represents a substantial improvement over clinician judgment alone for appropriate risk stratification. 44,45 However, it has some acknowledged limitations. For example, because age is the most heavily weighted variable in 10-year risk models derived from populations that span the adult age spectrum, in younger adults (< 45 years old in men and < 65 years old in women), modest elevations in risk factors have little effect on 10-year risk. 46,47 Even younger 26 adults with substantial risk factor burden may still have 10-year risk estimates well below 10%, although their remaining lifetime risks may exceed 50% on the basis of these risk factors. ⁴⁸ This is not a problem of the 10-year risk score per se, nor incorrect risk prediction, but rather a function of the decision thresholds that are imposed on the risk estimates and the short time horizon imposed by 10-year risk estimation.

The magnitude of risk factor levels that are needed to reach moderately high (10% to 20% 10-year predicted risk) and highallow more men and particularly more women in their 50s to guidelines recommend. 50 exceed 10% and 20% predicted 10-year risk. 47

adults to identify those with premature atherosclerosis.

SCREENING FOR CARDIOVASCULAR DISEASES

Screening for the presence of atherosclerotic coronary artery disease (CAD), or CVD in general, is potentially attractive on adequately investigated as yet.

Despite the immediate appeal of cardiovascular screening lesson that screening for vascular disease is not without hazards. with such common tests as coronary calcium measurement, 50 critical assessment of one of the most promising of the Evidence is strongest methods for early detection of CAD, coronary artery calcium (CAC) measured by rapid computed tomography, provides enlightening insights into the negatives of such screenings. On the one hand, recent reports from the Multi-Ethnic Study of Atherosclerosis (MESA) 30 convincingly demonstrate that ČAC testing is predictive of coronary events in whites, blacks, Hispanics, and Chinese Americans. Indeed, MESA investigated and found that increased CAC scores are predictive over and above traditional risk factors in both men and women. 30.55 Despite showing unusually strong hazard ratios for major coronary events ranging as high as approximately 7 for those with elevated CAC scores compared with a reference group with CAC = 0, MESA results also showed that CAC has a low positive predictive value for nearterm events regardless of the cutoff chosen for a positive test

result. 30 Depending on the cutoff chosen for a positive test result, sensitivity is also relatively poor. In MESA, for a cutoff value of CAC > 0 as the definition of a positive test result, sensitivity for near-term events was 91% but specificity was only 51% in this population of relatively low risk people (major coronary event rate during 3.9 years of median followup was slightly above 1%, with estimated 10-year risk extrapolated to approximately 5%). 42 With a cutoff of CAC > 0 for a positive test result, positive predictive value is only 2%. ⁵⁶ Furthermore, the positive predictive value would be substantially lower in younger populations than in the MESA cohort, which had a mean age of 62 years.

When a more stringent cutoff of CAC > 100 is used, sensitivity risk (> 20%) thresholds in the ATP III risk assessment for near-term major coronary events drops markedly to only 63%, algorithm has recently been investigated. Cavanaugh-Hussey and positive predictive value remains low at 3.5%. 56 Thus, although and colleagues 46 entered ranges of risk factor levels into the hazard ratios are impressive for CAC and far exceed those for ATP III risk assessment tool for men and women from the age carotid intima-media thickness, ^{37,51} CRP, ³⁸ or any of the traditional of 30 to 75 years. For almost all combinations of risk factors, risk factors for CHD, 30 positive predictive values for CAC even with extreme values, nonsmoking men < 45 years old and measurement are poor. Indeed, even with these impressive hazard essentially all women < 65 years old have 10-year predicted ratios, the C-statistic improves only from 0.79 for risk factors alone risks below 10%. Thus, many younger patients with significant as a predictor of CHD events to 0.83 for risk factors plus CAC score. risk factor burden do not reach treatment thresholds based on Thus, CAC scoring as a strategy to identify individuals at high risk current ATP III recommendations. A similar study ⁴⁷ has of near-term events across essentially the entire population, as has recently evaluated the updated Framing ham risk profile ⁴⁹ for been proposed by some, ⁵⁷ will likely not be feasible. It may, global CVD (not just CHD) risk, with somewhat similar however, still be very useful to reclassify risk among groups deemed findings, although the use of an expanded CVD endpoint does to be at intermediate risk by traditional risk models, as current

An important distinction between cancer screening tests and Several alternatives exist to address some of the limitations those available for cardiovascular risk assessment is availability (or of 10-year risk estimates, particularly for younger adults. One the lack thereof for CVD tests) of robust cost effectiveness or alternative is to change the horizon for risk estimation, to outcomes data from clinical trials of screening strategies provide long-term or lifetime risk estimates for CVD incorporating these modalities. We do not have a full understanding endpoints. Another possible solution would be to perform of the risks of CVD screening; nor do we have a full accounting of subclinical disease imaging in essentially all middle-aged benefits and costs. These issues have been the major cause for lack of endorsement of such tests as coronary calcium scoring for the general population by groups that include the USPSTF 54 and the UK National Health Service Research and Development Health SUBCLINICAL Technology Assessment Program. 58 Not only do we lack evidence that these sorts of screening confer a substantial benefit, we know that there is also a risk of harm. Harms of screening include false reassurance due to a test with low sensitivity, anxiety due to a falsepositive test result, and in the case of certain tests such as its face. However, it is fair to say that substantial further mammography and coronary computed tomography, radiation investigation of these screening strategies is warranted before exposures 59-61 as well as the additional anxiety and possible risks widespread adoption of these technologies is implemented. due to discovery and evaluation of incidental findings that result in Re-examination of Box 26-1 suggests that several of the World considerable added medical expense and invasive procedures. 62 The Health Organization criteria for screening have not been added burden of follow-up invasive angiography (that might or might not be necessary) observed after coronary calcium scoring is a

Radiographic mammography is recommended by the USPSTF, 63 B-mode ultrasound examination for intima-media thickness, 51 with a quality rating of B (ie, this level recommends that clinicians or other tests, 52 routine population-wide screening for provide this service to eligible patients on the basis of at least fair subclinical CAD or CVD has met with nearly universal evidence from clinical trials that the service improves important negative recommendations by expert panels to date. ^{29,53,54} A health outcomes and concludes that benefits outweigh harms).

26

for women aged 50 to 69 years, the age group generally included in screening trials. 63 With the identical review standards, the USPSTF against routine screening recommends electrocardiography, exercise treadmill testing, or rapid computed 9. tomography scanning for coronary calcium for either the presence of severe coronary artery stenosis or the prediction of CHD events in adults at low risk for CHD events (ie, screening of the general asymptomatic population). The USPSTF has concluded that there is at least fair evidence that these tests are ineffective or that harms 12. Vine DL, Hastings GE: Ischemic heart disease and cholesterol. Absolute risk more informative outweigh benefits. 54

measurement pertain to what is arguably the best and best-studied screening test currently available for detection of subclinical CAD. 14. Rembold CM: Number needed to screen: development of a statistic for disease screening. BMJ Potential concerns are magnified when applied to measures with large potential sources of error (relative to signal), such as carotid intima-media thickness and arterial stiffness measures. There are substantial difficulties associated with broad application of these 17. Law MR: The number needed to screen—an adaptation of the number needed to treat. J Med newer technologies in clinical practice, including technical issues (especially technician dependence of scanning), reader error and 18. Elmore JG, Barton MB, Moceri VM, et al: Ten year risk of false positive screening bias, and lack of agreement on the most appropriate parameters to be measured and the best instruments to use. It is also largely unclear 20. Ransohoff DF, Harris RP: Lessons from the mammography screening controversy: can we how often any of the screening tests should be repeated or what a meaningful change in the results may be.

CONCLUSION

Additional studies of screening strategies, including their impact on patient outcomes, are urgently needed to define the role of novel risk markers and subclinical disease imaging in screening for risk and subclinical CVD. These studies should be designed to answer the full 25. Pignone MP, Phillips CJ, Lannon CM: Screening for lipid disorders, systematic evidence range of issues and criteria needed before routine screening for the asymptomatic patient is recommended. The literature on screening tests is full of lessons indicating that we cannot assume clinical benefit, effectiveness, or cost-effectiveness simply because the tests 27. Genest J, McPherson R, Frohlich J, et al: 2009 Canadian Cardiovascular Society/Canadian we employ are feasible, have anatomical or physio logical face validity, or are strongly predictive of outcomes.

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CHAPTER 27

Role of Vascular Computed Tomography in Evaluation and Prevention of Cardiovascular Disease

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KEY POINTS

- Global risk assessment approaches for coronary heart disease, such as Framingham risk scores, underestimate longterm risks, especially in young men and postmenopausal women with multiple risk factors.
- Screening modalities, such as non-contrast-enhanced CT detection of coronary artery calcium, improve the ability to accurately predict risk in vulnerable groups and add information above and beyond global risk assessment.
- Absence of coronary artery calcification is associated with a very low risk of future CAD, significant stenosis, myocardial ischemia, and acute coronary syndrome during the next 5 years in nonsmokers.
- Guidelines reiterate that intermediate-risk individuals are the best patients for coronary artery calcium testing.
- Serial coronary artery calcium testing is currently not recommended because data for its prognostic significance are limited.
- Coronary artery calcium testing also appears to improve lifestyle changes and medication adherence.
- Coronary CTA has the potential to provide comprehensive information about the location, severity, and characteristics of

atherosclerotic plaque, especially noncalcified plaque; however, there are currently no recommendations for its use as a screening tool in asymptomatic persons.

 The prognostic value of CTA-based plaque subtypes and burden above and beyond coronary artery calcium scores is not yet established.

Cardiovascular disease (CVD) is the leading cause of mortality worldwide, with coronary artery disease (CAD) accounting for nearly half of all CVD deaths. 1-3 Although it is highly prevalent in the Western world, in the next 15 years, an estimated 25 million people will die of stroke or heart disease, with 80% of this burden in developing countries. ² Emerging data suggest that although the mortality rate after the occurrence of clinical heart disease myocardial infarction [MI]) significantly decreased during the last two decades, the incidence of new onset of CAD has remained relatively stable during this period. 4-6 This finding points to the fact that although we have made great strides in secondary prevention, we have failed in our primary efforts of decreasing the rate of newonset CAD.

In approximately half of the individuals, the initial presentation of CAD is either MI or sudden death. ¹ Unfortunately, conventional risk factor assessment predicts only 65% to 80% of future cardiovascular events, ¹ leaving many middle-aged and older individuals to manifest a major cardiovascular event despite being classified as low risk by the Framingham risk estimate. Because half of first major coronary events occur in asymptomatic individuals, clinicians who want to implement appropriate primary prevention therapy must be able to accurately identify at-risk individuals.

Clinical decision making for primary prevention of CAD in asymptomatic - individuals is traditionally guided by an initial estimate of the impact of single or clustered laboratory and physical factors as they relate to the risk of a coronary event. Preventive - strategies are then modified and implemented after economic (personal,

insurance provider, national impact) and individual (compliance, side effects) consequences of treatment versus treatment are taken into account. Recommendations for diet, weight loss, and exercise offer little or no risk to the patient and yield significant long-term benefits. 7 Most decisions for clinical (ie, pharmacologic) intervention, specifically those related to lipid lowering, are driven by perception of risk and achievement of goals for a given individual that are derived from large studies applied to both heterogeneous and homogeneous populations.

Screening for subclinical atherosclerosis in an effort to better identify persons at risk for CAD has been of increasing interest during the past decade. Cardiac computed tomography (CT) has shown promise in this regard, with a significant base of research and recent guidelines that have been established to support its use in selected groups of patients. Its use to better identify patients who might benefit from initiation or intensification of risk factor modification efforts is paramount to prevention efforts. This chapter details the current and future role of cardiac CT. The following discussion examines the methods, value, and future of cardiac CT in overall CAD in primary risk stratification and preventive strategies.

CURRENT CORONARY ARTERY DISEASE RISK STRATIFICATION GUIDELINES

A primary recommendation of the major advisory bodies is that all adults should undergo an office-based assessment as the

444 initial step to identify those at higher risk for CAD events. One ATHEROSCLEROSIS AND CORONARY ARTERY approach endorsed by both the American Heart Association (AHA) and the American College of Cardiology (ACC) and adopted by the National Cholesterol Education Program Considering that atherosclerosis is the major underlying culprit for (NCEP) Adult Treatment Panel III (ATP III) is to apply a modification of the risk prediction algorithm derived from the systolic blood pressure to estimate a person's 10-year risk for develop

I mention of a "hard" coronary heart disease (CHD) event (MI or CHD death). 8

Three levels of risk are defined on the basis of the probability of the occurrence of CHD in the next 10 years: < 10% (low), 27 10% to 20% (intermediate), and > 20% (high). 4 be offered general public health recommendations in the short term, and they can usually avoid further risk assessments for approximately 5 years. At the other end of the risk spectrum, highrisk patients are those with established CAD or other clinical forms of atherosclerotic disease, suffer from diabetes mellitus, or of exposure to the specific level of risk factors. 13 frequently are older patients with multiple other CAD risk factors reduce CAD-specific and total CVD risk. Finally, a sizable group of individuals fall into the intermediate -risk category. Patients in this group do not currently qualify for the most intensive risk factor interventions, yet they have one or more risk factors that exceed desirable levels or multiple borderline risk factors. Often, there is not a definite need to begin or to intensify therapy; however, many such persons do have underlying subclinical atherosclerosis that would place them at higher future risk of CAD stratification testing, if it is feasible, practical, targeted, and behavioral changes. 8

Determination of the intensity of treatment for primary prevention by use of global risk assessment algorithms such as the Framingham risk score (FRS) system employs data from volumetric acquisitions. population-based studies and does not take into account an individual's actual burden of atherosclerotic disease, which is believed to be the main culprit in the development of clinical CAD. Indeed, such assessments may fall short if they are solely relied on for management decisions for the individual patient as these risk estimates are significantly age dependent; CORONARY as a result, there is a great tendency to underestimate risk in **ASSESSMENT OF CORONARY** younger individuals, who may be more appropriate ARTERY CALCIFICATION candidates for early initiation of aggressive preventive women younger than 70 years are considered low risk. 10

NCEP guidelines to underestimate the risk for CAD was even hearts. more pronounced in women, with only 18% of women qualifying for pharmacotherapy for primary prevention; 58% of these patients had low-density lipoprotein cholesterol (LDL-C) concentration < 130 mg/dL, and 40% had LDL-C < 100 mg/dL. 11

DISEASE

the development of clinical CAD, detection of individuals with subclinical atherosclerosis may aid in supplementing current global Framingham Heart Study that incorporates a patient's age, risk assessment approaches by more clearly identifying high-risk total cholesterol concentration, high-density lipoprotein individuals who harbor advanced preclinical atherosclerosis. cholesterol (HDL-C) concentration, smoking status, and Screening studies to detect occult cancers, such as breast and colon cancer, are recommended in appropriate-risk adults to help prevent life- threatening conditions. Although atherosclerotic vascular disease accounts for more death and disability than all types of cancer, a screening tool to detect significant subclinical atherosclerosis and to target prevention of future cardiovascular events has not yet been adopted. 12

A minority of patients with CAD do not exhibit traditional risk Individuals with CAD risk < 10% are at presumed low risk of factors, such as hypertension, elevated cholesterol, obesity, and coronary events and can be reassured about their risk status smoking. In addition, many patients with such risk factors do not without further risk assessment testing. Those with low risk are to develop CVD. Furthermore, there is substantial variation in the severity of CAD at every level of risk factor exposure. This variation in disease is probably due to a number of factors, including genetic susceptibility, presence of intrinsic biochemical and extrinsic environmental risk factors that are yet to be identified, and duration

Established noninvasive methods of evaluating CAD, such as in accordance with NCEP ATP III guidelines; high-risk stress testing, generally identify only patients with advanced asymptomatic people should have all CAD risk factors treated to atherosclerotic disease leading to a flow-limiting coronary stenosis and myocardial ischemia. The long-term risk of CAD, however, is more closely related to atherosclerosis plaque burden and stability than to the extent of a particular stenosis. ¹⁴ There is growing interest in quantifying and characterizing atherosclerosis in its preclinical, pre flow-limiting phase so that appropriate preventive strategies can be instituted before an adverse event occurs.

With understanding of the need for noninvasive tests to assess atherosclerotic plaque, cardiac CT has evolved rapidly, with than global risk assessment would predict. Intermediate-risk increasing ability to visualize the amount of coronary artery patients are the most likely to benefit from further risk calcium. 15 Cardiac CT has been challenging, given rapid cardiac motion, small vessel diameters, tortuous anatomic patterns, and effective at further defining risk or in motivating effective overlapping cardiac structures. Current 64-slice multirow detector computed tomography (MDCT) systems have faster gantry rotation speed, resulting in better temporal resolution and the better z- axis spatial resolution made possible by thin collimations with extensive

NON-CONTRAST-ENHANCED COMPUTED

TOMOGRAPHY:

strategies to reduce risk for development of clinical CAD. 9 In During the past decade, there has been marked increased interest in addition, the risk of a future CVD event in women is likely to the clinical use of cardiac CT scanning to identify and quantify the be underestimated by these approaches, and nearly 90% of amount of coronary artery calcified plaque (CAC). Calcification of the atherosclerotic plaque occurs by an active process of This point can be illustrated in a study by Akosah and mineralization with deposition of hydroxyapatite crystals and not colleagues, 11 in which the authors assessed a simple question: simple mineral precipitation. It begins in the very early stages of How do NCEP guidelines classify young men and women atherosclerosis. Studies have demonstrated that electron beam presenting with an acute MI as their first manifestation of tomography (EBT) is a highly reliable method for identification of CAD? The study findings demonstrated that among adults arterial calcification with a high sensitivity for detection of aged < 65 years with an acute MI, only 25% of patients would significant atherosclerosis. Rumberger and colleagues 16 have have met ATP III criteria, which are based on the FRS, for demonstrated a strong relationship (r = 0.90) between CAC pharmacotherapy at the time of admission. The tendency of measured by EBT and direct histological plaque areas in autopsy

compensation for increasing area of mural plaque.

and luminal stenosis, coronary calcium scores calculated by EBT tool for the identification of patients at risk.

METHODS OF ASSESSING CORONARY ARTERY **CALCIUM**

Modalities for Calcium Coronary Artery **Determination**

In the near past, CAC had generally been assessed by EBT; however, with a rapid explosion of use of MDCT in recent years. This has been a widely used modality to assess the extent and severity of underlying coronary calcification. 19 Neither modality requires intravenous administration of contrast material to determine CAC. In general, EBT uses a unique technology enabling ultrafast scan acquisition times in the high-resolution, single-slice mode with continuous, nonoverlapping slices of 3-mm thickness and an acquisition time of 100 msec/tomogram in a prospective manner. Electrocardiographic (ECG) triggering is done during end systole or early diastole at a time determined from the continuous ECG tracing recorded during scanning. Historically, the most common trigger time used is 80% of the RR interval. However, this trigger occurs on or near the P wave during atrial systole, and the least cardiac motion among all heart rates occurs at 40% to 60% of the RR interval. 19,20

The current generation of MDCT systems is capable of acquiring up to 320 sections of the heart simultaneously with ECG gating in either a prospective or retrospective mode. These MDCT systems have two main modes of scanning, which depend on whether the patient on the CT couch is advanced in a stepwise fashion (axial, sequential, or conventional mode) or continuously moved at a fixed speed relative to the gantry rotation (helical or spiral mode). Coronary calcification is determined in axial mode with the use of prospective ECG triggering at predetermined offset from the ECGdetected R wave. With prospective gating, the temporal resolution of the MDCT system is proportional to the gantry speed, which determines the time to complete one 360-degree rotation.

For the reconstruction of each slice, data from a minimum of 180 degrees plus the angle of the fan beam are required (approximately 220 degrees of the total 360-degree rotation). The most commonly used 64-slice scanners have rotation gantry speeds up to 330 msec. ¹⁹ MDCT imaging protocols vary among different camera systems and manufacturers. In general, 40 consecutive 2.5- to 3-mm-thick images are acquired per cardiac study. Calcified lesions are defined as two or three adjacent pixels with a tomographic density of > 130 HU. Effective pixel size for a reconstruction matrix of 512 x 512 pixels with a common field of view of 26 cm is 0.26 mm².

Measurement of Coronary Artery Calcium Burden

On non-contrast-enhanced cardiac CT, CAC is generally defined as a hyperattenuated lesion above a threshold of 130 HU with an area of three adjacent pixels (at least 1 mm²). There are currently two CT calcium scoring systems widely used, the original Agatston method ¹⁹ and the "volume" score method. ²⁰ The Agatston method involves

Although the total atherosclerotic plaque burden was associated multiplication of the calcium area by a number related to CT density strongly with the total calcium burden, not all plaques were found and, in the presence of partial volume artifacts, can be variable. With to be calcified. Moreover, within a given coronary artery, there is a this method, the area for all pixels above a threshold of 130 HU is poor correlation and wide variation between the degree of plaque calculated at every 3-mm slice and multiplied by a density factor. 19 calcification and extent of luminal stenosis on coronary Partial volume effects lead to higher peak values for small lesions angiography. 17,18 This may be due, at least in part, to individual (but not for large ones). On the other hand, the volume method variations in coronary artery remodeling, whereby the luminal developed by Callister and associates 20 appears to somewhat cross-sectional area or external vessel dimensions enlarge in resolve the issue of slice thickness and spacing by computing a volume above threshold. As a result, it appears to be less dependent Despite the lack of a site-by-site correlation between calcification on minor changes in slice thickness. However, our group has previously demonstrated in nearly 10,000 patients that there give a close approximation of the total atherosclerotic burden. 18 appeared to be an excellent correlation between the scoring Because research has shown that burden of disease and methods, and they show similar characterization when applied cardiovascular risk are accounted for by more than focal luminal properly. 21 Both methods calculate lesion-specific scores within the 27 stenosis, non-contrast-enhanced cardiac CT is a potentially powerful left main, left circumflex, left anterior descending, and right

left main, left circumflex, left anterior descending, and right coronary arteries and provide total scores for each artery and a total sum across all four arteries. An example of significant coronary calcium is shown in Figure 27-1.

MULTIDETECTOR COMPUTED TOMOGRAPHY
COMPARED WITH ELECTRON BEAM
TOMOGRAPHY FOR DETECTION OF CORONARY
ARTERY CALCIUM

The comparability of MDCT- and EBT-derived CAC scores has been extensively explored. 22-24 The MDCT protocols vary considerably in these studies, ranging from conventional CT to single-slice CT (with either retrospective or prospective gating) to MDCT. EBT imaging was performed with the standard protocol conventionally used in was performed with the standard protocol conventionally used in routine clinical practice. Coronary calcification was defined as > 130
HU for EBT but varied from 90 to 130 HU for MDCT. Although high
correlation coefficients were reported between EBT and MDCT CAC
scores, there was significant variability in individual CAC scores
(range, 17% to 84%). In general, the interscan agreement for the
presence of CAC between EBT and 64-slice routine clinical practice. Coronary calcification was defined as > 130

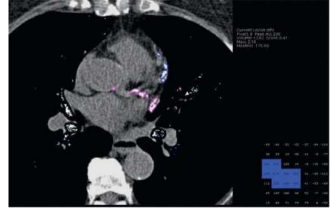


FIGURE 27-1 Example of significant coronary artery calcium from a multidetector CT scanner

446 MDCT is excellent (99%). There was a significant linear relationship between the scores from the two scanners, and the interscanner variability risk with the presence of CAC. In one of the largest observational between EBT and 64-slice MDCT was not significantly different. Bland-trials to date, Shaw and colleagues 29 reported all-cause mortality Altman analysis demonstrated a mean difference in scores of 8.3% by among 10,377 asymptomatic patients (4191 women and 6186 men) Agatston and 7.8% by volumetric calcium scoring. Compared with EBT, who had a baseline EBT study and were then observed for 5.0 ± 3.5 there were larger and more prevalent motion artifacts and larger mean years (Fig. 27-2). Most subjects had cardiac risk factors including a Hounsfield units with 64-slice MDCT (P< 0.001). At CAC scanning, 64- family history of CAD (69%), hyperlipidemia (62%), hypertension slice MDCT and EBT were comparable in Agatston and volumetric (44%), and current cigarette smoking (40%). The CAC score was a scoring. The interscan variability I between scanners is similar to the strong independent predictor of mortality, with 43% additional interscan variability of two calcium scores done on the same equipment. predictive value contained within the CAC score beyond risk factors However, heart rate control was achieved for this study for calcium scores. alone. Mortality significantly increased with increasing CAC score, Whether these results are repeatable without heart rate 27 control needs within men and women separately as well as within each to be further assessed.

CLINICAL VALUE OF CORONARY ARTERY CALCIFICATION IN ASYMPTOMATIC INDIVIDUALS

Efforts have been made to develop noninvasive diagnostic tools to help determine the extent of atherosclerosis in asymptomatic patients and to improve detection of those who would benefit from more intensive preventive thera pies, such as lipid-lowering medication and aspirin. This potential of coronary CT in the risk assessment protocol and management strategy is in accordance with the current philosophy of the NCEP and other organizations that stress the importance of matching therapy to the level of assessed risks. However, to establish the role of CAC testing in primary preventive strategies, important questions need to be answered. Is the information gained from coronary CT additive to assessments made by cheaper office-based estimations of risk? If so, which populations of patients are expected to benefit from testing?

Does Coronary Artery Calcium Independently Predict Coronary Artery Disease Events?

cardiovascular events is related to the total atherosclerotic plaque CAC and incremental sex-specific thirds of detectable CAC; these burden. 13,18 Although controversy exists as to whether calcified or rates were, respectively, 0.4, 1.5, 4.8, and 8.7 (trend P < 0.0001) for noncalcified plagues are more prone to rupture, 14 extensive men and 0.7, 2.3, 3.1, and 6.3 (trend P < 0.02) for women. The calcification indicates the presence of both plaque morphologies. 16- association between CAC and CAD events remained significant 18 There is a direct relationship between the CAC severity and the after adjustment for CAD risk factors. Of note, in the largest single extent of atherosclerotic plaque; thus, the CAC score could be useful cohort study, Budoff and colleagues 35 showed risk-adjusted hazard for risk assessment of asymptomatic individuals and potentially ratios for total mortality ranging from 2.2 to 12.5 for CAC score guide therapeutics.

the prognostic value of CAC burden among asymptomatic individuals. ²⁵⁻³⁸ In general, there appears to be a consensus among CAC was determined by MDCT in 924 patients (443 men, 481 all studies that CAC is an independent predictor of CAD adverse women, aged 59.4 ± 18.7 years). During the 3-year follow-up period, outcome as well as of all-cause mortality after traditional risk factors the event rates for coronary revascularization, MI, and cardiac death are taken into account. Among 1172 asymptomatic patients in patients with volume scores above the 75th percentile were observed for 3.6 years after an initial EBT screening, no events significantly higher compared with the total study group, and no occurred in patients with a normal study, and the negative cardio vascular events occurred in patients with scores of zero. predictive value was 99.8% in patients with a CAC score < 100. Receiver operating characteristic (ROC) analysis demonstrated that These results showed a 5%, 7%, and 13% hard cardiac event rate in it outperformed both PROCAM and Framingham individuals with a CAC score > 80, > 160, and > 600, respectively. 25 The CAC score remained the best single predictor of risk after adjustment. Wong and colleagues 26 also showed that the CAC score severity predicted subsequent cardiovascular events independent of age, gender, and patient risk factor profile. Raggi and coworkers 27 studied more than 600 asymptomatic patients who were referred for EBT screening and then observed for 32 ± 7 months. Both the absolute CAC score and the relative CAC score percentiles adjusted for age and gender predicted subsequent death and nonfatal MI. Hard cardiac events occurred in only 0.3% of subjects with a normal EBT study, but this increased to 13% in those with a CAC score > 400. A very high CAC score > 1000 may portend a particularly high risk of death or MI (25% per year) in individuals who are not treated with standard secondary prevention measures.

Larger studies have reported an approximately 10-fold increased Framingham risk group (low-, intermediate-, and high-risk persons). 29 In addition, in the South Bay Heart Watch study, 1196 asymptomatic patients were observed (median = 7.0 years), and it was demonstrated that the CAC score added predictive power beyond that of standard coronary risk factors and C-reactive protein ³⁰ (see Table 27-1). The results of the St. Francis Heart Study, which is a prospective registry of 5585 asymptomatic individuals, mirrored previous retrospective studies and confirmed the higher event rates associated with increasing CAC scores. ³¹ CAC scores > 100 were associated with relative risks of 12 to 32, thus achieving secondary prevention equivalent event rates > 2%/year. 31 The Rotterdam Heart Study ³² investigated 1795 asymptomatic participants (mean age, 71 years) who had CAC and measured risk factors. During a mean follow-up of 3.3 years, the multi variate-adjusted relative risk of coronary events was 3.1 for calcium scores of 101 to 400, 4.6 for calcium scores of 401 to 1000, and 8.3 for calcium scores > 1000. Similarly, in a younger cohort of asymptomatic persons, the 3-year mean follow-up in 2000 participants (mean age, 43 years) showed that coronary calcium was associated with an 11.8-fold increased risk for incident CHD (CAD) (P < 0.002) in a Cox model controlling for the FRS. 33

The Cooper Clinic Study ³⁴ included more than 10,000 adults who were 22 to 96 years of age and free of known CAD. During a mean follow-up of 3.5 years, 81 hard events (CAD death, nonfatal MI) occurred. Age-adjusted rates (per 1000 person-years) of hard events The likelihood of plaque rupture and the development of acute were computed according to four CAC categories: no detectable categories of 11-100 to > 1000 relative to 0, with CAC scores Table 27-1 summarizes the findings of all major studies assessing providing significant incremental information about risk factors. Finally, in a German study by Becker and coworkers, 36 the extent of

TABLE 27-1	Summary of Outcomes Studies with	Coronary Artery	/ Calcification in Asymptomatic I	ndividuals
Author (Year)	Type of Study and	Follow-up (Number of Events	Results
Arad et al ≈ (2000)	Population Observational study, referral	years) 3.6	15 nonfatal MI, 21	OR of 20 for CAC scores > 160 compared with those with CAC
	based N = 1172; age, 53 ± 11 years		revascularizations, 3 deaths	scores < 160
Wong et al 26 (2000)	Observational study, referral based N = 926; mean age, 54 years	3.3	6 nonfatal MI, 20 revascularizations, 2 CVAs	Overall, patients with CAC score > 271 had a risk ratio of 9 for a CHD event.
Raggi et al 27 (2001)	Observational referral based study N = 676; mean age, 52 years	2.7	21 nonfatal MI, 9 deaths	CAC score was predictive of hard CAD events, with an OR of 22 for the CAC score 90% percentile.
Kondos et al 28 (2003)	Observational study, referral based N = 5635; age, 30-76 yr; 26% women	3.1	37 nonfatal MI, 166 revascularizations, 21 deaths	RR of 124 in men with soft events in the highest quartile (CAC 170-7000) Higher CAC scores added incremental prognostic information to conventional CAD risk assessment in men for hard CHD events.
Shaw et al 20 (2003)	Observation data series, referral based N = 10,377; age, 30-85 years	5	249 all-cause mortality	CAC score an independent predictor of mortality with RR 4.0 for score of 401-1000
Greenland et al » (2004)	Prospective population based study N = 1312; age, > 45 years	7	68 nonfatal MI, 16 deaths	HR of 3.9 for CAC score > 301 CAC score able to modify predicted risk obtained from FRS alone (0.73 for FRS alone and 0.78 for FRS and CAC combined)
Arad et al 31 (2005)	Prospective population based study N = 4613; age, 50 to 70 years	4.3	40 nonfatal MI, 59 revascularizations, 7 CVAs	RR for CAD events with CAC > 100 was 11. Overall, it was superior to FRS in prediction of events (ROC curve of 0.79 versus 0.69; <i>P</i> = 0.006).
Vliegenthart et al 22 (2005)	Prospective population based study N = 1795; age, 62 to 85 years	3.3	40 nonfatal MI, 11 revascularizations, 38 CVAs	Compared with those with CAC < 100, the RR for events was 3.1, 4.6, and 8.3 for CAC 101-400, 401-1000, and > 1000, respectively. There was a statistically significant high relative risk, > 8, for those with CAC scores > 1000 regardless of Framingham 10-year risk score < 20% versus > 20%.
Taylor et al »(2005)	Prospective cohort study 1627 men, 356 women; age, 40 to 50 yrs (Army based)	3	9 ACS events	2% of men with CAC had events versus 0.2% without CAC (P < 0.0001). Controlling for FRS, presence of CAC was associated with an independent 12-fold increase in relative risk. No events in women
LaMonte et al 34 (2005)	Retrospective study 6835 men, 3911 women; age, 22-96 years	3.5	81 MI/CAD death, 206 revascularizations	Age-adjusted rates per 1000 person-years were computed according to 4 CAC categories: 0 CAC and incremental sexspecific thirds of detectable CAC. The rates were 0.4, 1.5, 4.8, and 8.7 for men and 0.7, 2.3, 3.1, and 6.3 for women.
Anand et al » (2006)	Prospective study 510 asymptomatic type 2 diabetic subjects; age, 53 ± 8 yrs	2.2	22 (2 coronary deaths, 9 nonfatal MI, 3 ACS, 3 CVA, and 3 late revascularizations)	The overall rate of death or MI by CAC categories ($<$ 100, 101-400, 401-1000, and $>$ 1000) was 0 (n = 0), 2.6 (n = 2), 13.3 (n = 4), and 17.9% (n = 5), respectively ($P <$ 0.0001).
Budoff et al »(2007)	Observation data series, referral- based N = 25,253; mean age, 65 ± 11 years	6.8	510 all-cause deaths	Compared with those without CAC, the risk-adjusted relative risk ratios for CAC were 2.2-, 4.5-, 6.4-, 9.2-, 10.4-, and 12.5-fold for scores of 11 to 100, 101 to 299, 300 to 399, 400 to 699, 700 to 999, and > 1000, respectively (<i>P</i> < 0.0001).
Detrano et al ³⁰ (2008)	Prospective population based study	3.4	162 CHD events (72 myocardial infarctions, 17 CHD deaths, 73 revascularizations)	Overall, the FRS-adjusted risk was 28% higher with doubling of CAC scores. CAC was equally predictive in all ethnic groups.
Becker et al ³⁸ (2008)	Prospective population based study	3.3	179 (65 cardiac deaths, 114 MI)	CAC score > 75th percentile was associated with a significantly higher annualized event rate for MI (3.6% versus 1.6%; P < 0.05). No cardiac events were observed in patients with CAC = 0.

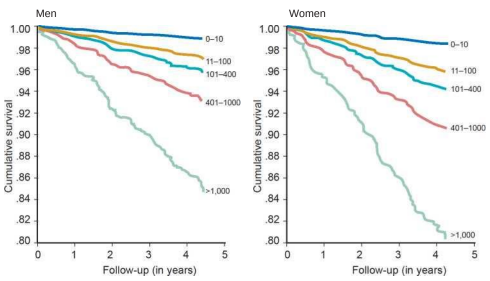


FIGURE 27-2 Risk of total mortality by calcium category in 10,377 asymptomatic individuals. (From Shaw LJ, Raggi P, Schisterman E, et al: Prognostic value of cardiac risk factors and coronary artery calcium screening for all-cause mortality. Radiology 228:826, 2003.)

				Even	ts / N						
Summary RR Ratio	CACS	RR	(95% CI)	Higher Risk	Low Risk*	P	0.01	0.1	1	10	100
Average risk	1-112	1.9	(1.3-2.8)	67/9,514	45/12,163	0.001					
Moderate risk	100-400	4.3	(3.1-6.1)	110/5,209	49/11,817	< 0.000					
High risk	400-999	7.2	(5.2-9.9)	182/3,940	49/8,649	< 0.000					
Very high risk	1,000	10.8	(4.2-27.7)	14/196	6/905	< 0.000					
, 0			,				0.01	0.1	1	10	100
							Lower ri	۔۔۔ ہے باہ		\ Hic	shor rick

FIGURE 27-3 Meta-analysis of relationship of CAC scores (CACS) with CHD outcomes. (From Greenland P Bonow RO, Brundage BH, et al: ACCF/AHA 2007 clinical expert consensus document on coronary artery calcium scoring by computed tomography in global cardiovascular risk assessment and in evaluation of patients with chest pain: a report of the American College of Cardiology Foundation Clinical Expert Consensus Task Force [ACCF/AHA Writing Committee to Update the 2000 Expert Consensus Document on Electron Beam Computed Tomography] developed in collaboration with the Society of Atherosclerosis Imaging and Prevention and the Society of Cardiovascular Computed Tomography. J Am Coll Cardiol 49:378, 2007. Reproduced with permission.)

models (P < 0.0001), in which 36% and 34% of MIs occurred in the high-risk cohorts, respectively.

The utility of CAC testing was also recently described in CAD-equivalent individuals, that is, those with diabetes mellitus . Risk factors and CAC scores were prospectively measured in 510 asymptomatic type 2 diabetic subjects (mean age, 53 ± 8 years; 61% men) without prior CVD, with a median follow-up of 2.2 years. ³⁷ In the multivariable model, the CAC score and extent of myocardial ischemia were the only independent predictors of outcome. ROC analysis demonstrated that CAC predicted cardiovascular events with the best area under the curve (0.92), significantly better than the United Kingdom Prospective Diabetes Study risk score (0.74) and Framingham score (0.60). The relative risk to predict a cardiovascular event for a CAC score of 101 to 400 was 10.13, and it increased to 58.05 for scores > 1000 (P < 0.0001). No cardiac events or perfusion abnormalities occurred in subjects with CAC < 10 Agatston units up to 2 years of follow-up.

These findings were nicely summarized by the AHA/ACC expert consensus document on coronary artery calcium scoring, which took into account many of these studies. Compared with patients with no detectable coronary calcium, the relative risk ratio for CAC 100-400, 401-1000, and > 1000 was 4.3 (95% CI, 3.5-5.2; P < 0.0001), 7.2 (95% CI, 5.2-9.9; P < 0.0001), and 10.8 (95% CI, 4.2-27.7; P < 0.0001), respectively . Importantly, patients with CAC score of zero have a very low rate of CAD death or MI (0.4%) during 3 to 5 years of observation ³⁸ (Fig. 27-3).

At the same time, critics tend to point to limitations such as

potential study generalizability of self-referral cohorts, validity of the risk factor measures and resulting multivariable models used in the studies, and risk of test-induced bias. However, these concerns have been addressed by a report from the Multi-Ethnic Study of Atherosclerosis (MESA), a population-based cohort, which reported the utility of CAC scores in predicting future events. According to Detrano and coworkers, 39 among nearly 6800 asymptomatic individuals observed for a median of 41 months, the hazard ratio for future hard CAD events (MI or MIrelated death) with CAC 1-100 versus CAC = 0 was 5.3 (95% CI = 2.4-11.7; P < 0.0001). The respective hazard ratios with CAC 101-300 and > 300 were 10.8 (4.8-24.2; P < 0.0001) and 12.0 (5.4-26.5; P < 0.0001), with a 5-year cumulative incidence of CAD events directly associated with higher CAC scores, exceeding 10% in those with scores > 300 (Fig. 27-4). These risk ratios are very similar to those of published studies and confirm the pooled summary findings previously reported and lay to rest any concern about the prognostic value of CAC testing. 38

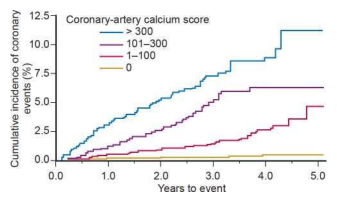


FIGURE 27-4 Cumulative incidence of CAD events according to increasing CAC score categories in the Multi-Ethnic Study of Atherosclerosis. (From Detrano R, Guerci AD, Carr JJ, et al: Coronary calcium as a predictor of coronary events in four racial or ethnic groups. N Engl J Med 358:1336, 2008.)

What Is the Value of Testing for Coronary Artery Calcium Across Ethnic and Racial Groups?

Most of the published data to date have related to white populations; however, two studies have addressed the value of CAC in other ethnic groups. First, Nasir and coworkers, 40 in nearly 15,000 ethnically diverse self-referred patients, assessed the role of CAC for the prediction of all-cause mortality. In comparison of prognosis by CAC scores in ethnic minorities and non-Hispanic whites, relative risk ratios were highest for African Americans, with scores > 400 exceeding $16.\overline{1}$ (P < 0.0001). Hispanics with CAC scores > 400 had relative risk ratios from 7.9 to 9.0; Asians with CAC scores > 1000 had relative risk ratios 6.6fold higher than those of non-Hispanic whites (P < 0.0001). Second, the utility of CAC testing has also been reported in the prospective MESA study. 39 According to Detrano and coworkers, the risk associated with a doubling of the CAC score (a 1-unit increase in log 2 [CAC + 1]) for a hard CAD event was 1.3 (1.2-1.4) in whites, 1.5 (1.3-1.7) in African Americans, 1.3 (1.1-1.5) in Hispanics, and 1.4 (1.1-1.8) in Chinese. These findings firmly establish that CAC scores provide significant information in all four major ethnic groups in the United States.

Does Coronary Artery Calcium Add Incremental Value to Global **Risk Estimates?**

The extent of CAC has been shown in several studies to predict cardiac events in symptomatic and asymptomatic individuals. However, decisions about the predictive utility of new tests should be based on the additional utility of a new test for risk prediction. The most important question about use for primary CAD risk stratification is whether it is predictive above and beyond the current standard risk assessment method of choice, the FRS, which is an inexpensive, easily available, and officebased tool. One way to determine the additive utility of a new test is through the use of ROC curve analyses. The ROC curve is a plot of true-positive rate versus false-positive rate over the entire range of possible cutoff values. The area under the ROC curve (AUC) ranges between 1.0 for the perfect test and 0.5 for the useless test.

Studies comparing predictive capacity of conventional and newer biomarkers for prediction of cardiovascular events consistently demonstrated that adding a number of newer bio markers (such as C-reactive protein, interleukins, and other proposed risk stratifiers) changes the C-statistic by only 0.009 (P = 0.08). Small changes such as these in the C-statistic suggest limited or modest improvement in risk discrimination with additional risk markers. However, CAC scanning has been shown to markedly improve the C-statistic, suggesting robust improvement in risk discrimination (Table 27-2).

Outcomes with Use of Coronary Artery Calcium Compared with Traditional Risk Factors				
Study	C-Statistic with Risk Factors, FRS	C-Statistic with Risk Factors, FRS plus CAC	<i>P</i> Value	
Arad et al ³¹ (St. Francis Heart Study)	0.69	0.79	0.0006	
Budoff et alas	0.611	0.813	<0.0001	
Anand et als7	0.60	0.92	<0.0001	
Becker et als	0.68	0.77	<0.01	
Detrano et al ³⁰ (MESA)	0.77	0.82	<0.001	

Raggi and colleagues 27 were among the first to assess the added contribution of CAC over and above the FRS. In a study of more than 10,000 asymptomatic individuals observed nearly 5 years, the C-statistic (from ROC curve analyses) for FRS in estimating risk of all-cause death was 0.67 (95% CI, 0.62-0.72; P < 0.0001) for women and 0.68 (95% CI, 0.64-0.73; P < 0.0001) for men. When CAC was added to this analysis, the C-statistic increased to 0.75 (95% CI, 0.70-0.80) for women (P < 0.0001) and 0.72 (95% CI, 0.68-0.77) for men (P < 0.0001), indicating asignificant improvement in mortality prediction. ²⁷

In a similar fashion, Greenland and coworkers 30 found that the ROC curve for prediction of CAD death or nonfatal MI was 0.68 for FRS plus CAC, which was significantly greater than that of the FRS alone (0.63; P < 0.001), with increasing levels of CAC associated with greater risk within each FRS group. Importantly, those in the intermediate-risk FRS group with high CAC scores had event rates as high as or higher than those of persons within the high-risk FRS group with lower CAC scores (Fig. 27-5). The recent population-based St. Francis Heart Study of 5585 asymptomatic individuals confirmed the findings of previous reports. The CAC score predicted CAD events independently of standard risk factors and C-reactive protein (P = 0.004), was superior to the FRS in the prediction of events (area under ROC curve of 0.79 ± 0.03 versus 0.69 ± 0.03 ; P = 0.0006), and enhanced stratification of those falling into the Framingham categories of low, intermediate, and high risk (P < 0.0001). ³¹ Similarly, an improvement in AUC from 0.77 to 0.82 was noted in the landmark MESA study. 39

More recently, Becker and coworkers ³⁶ demonstrated that among 1726 asymptomatic individuals observed for a median of 40 months, the area under the ROC curve for CAC scores (0.81; 95% CI, 0.78-0.84) was significantly larger than that for the FRS risk (0.63; 95% CI, 0.59-0.65), PROCAM (0.65; 95% CI, 0.6-0.68), and European Society of Cardiology scores (0.66; 95% CI, 0.62-0.6) (P = 0.03). Most recently, Polonsky and colleagues reported in the Multiethnic Study of Atherosclerosis that the addition of CAC to models with age, gender, and risk factors resulted in a net reclassification of $0.\overline{25}$ ($\overline{P} < 0.001$), where 23% of subjects with events were reclassi fied as high risk and 13% of those without events were reclassified as low risk. 40a In addition, in the Heinz Nixdorf Recall Study of 4129 subjects aged 45-75 without CHD at baseline, the addition of CAC to FRS not only improved the area under the curve from 0.68 to 0.75 (P < 0.003) but also resulted in a

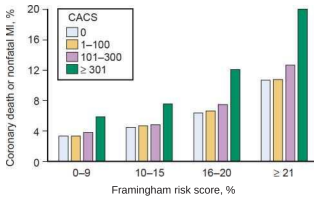


FIGURE 27-5 Predicted 7-year CHD event rates by CAC scores (CACS) within Framingham risk score groups. (From Greenland P, LaBree L, Azen SP, et al: Coronary artery calcium score combined with Framingham score for risk prediction in asymptomatic individuals. JAMA 291:210, 2004. Reproduced with permission.)

these findings not only support the contention that established associated with excellent survival, with all-cause mortality rates of cardiac risk factors possess a limited ability to quantify CAD risk 0.87 per 1000 per son-years (< 1% 10-year risk, or < 0.1% per year). but also provide evidence that CAC may add unique information In a similar fashion, Budoff and colleagues, 43 in a multiethnic to risk assessment. As we go forward, it will be important to prospective cohort, demonstrated that those individuals with document the degree to which bio markers or imaging tests, calcium scores of zero had a remarkably low rate of cardiovascular including CAC screening, can appropriately reclassify individuals events. In a median of 4.1 years of follow-up, the event rate for hard from standard risk factor assessment for the purposes of more CAD with CAC = 0 was 0.74 per 1000 participants (95% CI, 0.40accurately targeting the intensity of treatment.

ABSENCE OF CORONARY **ARTERY CALCIFICATION**

There is increasing interest in the absence of CAC (a calcium score of zero) as a "negative" cardiovascular risk factor. Absence of CAC may reliably exclude obstructive coronary disease in asymptomatic and selected symptomatic individuals and appears to be associated with a low cardiovascular event rate, suggesting that less aggressive pharmacotherapy may be indicated in this population. However, published event rates for individuals with zero CAC vary, probably because of differences in baseline risk, follow-up period, and outcome ascertainment and verification. Our group has examined the literature for the relevance of the absence of CAC in the context of three major categories: (1) its prognostic utility in categorizing both asymptomatic and symptomatic patients according to their risk for adverse events, (2) its relationship with the presence or absence of significant coronary artery stenosis by invasive coronary angiography, and (3) the degree of myocardial ischemia detected in those with the absence of CAC.

Prognostic Value of Absence of Coronary Artery Calcification in Asymptomatic and Symptomatic **Individuals**

In our meta-analysis, ⁴¹ outcomes consisting of 71,595 asymptomatic *individuals*, 29,312 patients (41%) did not have any evidence of CAC. In a pooled follow-up of 50 months, only 0.47% of asymptomatic we do stress that individuals who smoke and have diabetes, even individuals without CAC (154 of 29,312) suffered a cardiovascular in the absence of CAC, should be treated according to existing event during follow-up compared with 4.14% with CAC (1749 of guidelines. 42,283). In a similar fashion, the meta-analysis revealed that among 3924 symptomatic patients (60% male), 921 patients (23%) did not have any evidence of CAC. In these symptomatic individuals, only 17 of 921 patients (1.8%) without CAC suffered a cardiovascular Myocardial Ischemia in the Absence of Coronary event during follow-up of 42 months compared with 270 of 3003 patients (9.0%) with CAC.

Our group has further explored the prognostic value of absence of CAC in two different cohorts. Blaha and associates 42 showed that in a cohort of 44,052 asymptomatic middle- aged patients free of

significant net reclassification of 21.7% of patients. 40b Importantly, known CAD observed for up to 13 years, a CAC score of zero was 1.37); whereas for all CAD, those with CAC = 0 had 1.25 (95% CI, 0.78-2.02) events per 1000 person-years at risk.

However, even in the absence of CAC, relatively more events occur among those with higher risk, especially in diabetics and smokers. 42,43 The potential mechanisms may include the presence of underlying noncalcified components, rapid development of atherosclerosis, and plaque destabilization . Whereas the relative risk of events is higher in the presence of low CAC, the absolute event rate remains low. In an appropriately selected non-high-risk patient, the absence of CAC could potentially be used as a rationale to emphasize lifestyle therapy, to scale back on costly preventive pharmacotherapy, and to refrain from frequent cardiac imaging

Given the low 1% 10-year risk in this population, a drug that produces a 30% relative risk reduction would have to be given to more than 300 patients for 10 years to prevent one death (number needed to treat, ~333 for 10 years). 42 Although current guidelines do not recommend that preventive therapies such as lipidlowering medications be downregulated in the absence of CAC, emerging data 41-43 suggest that aggressive management in this cohort is not warranted if one does not qualify according to NCEP guidelines. As such, physicians may consider emphasizing appropriate lifestyle therapy, using less pharmacotherapy, and not ordering costly cardiac imaging studies if there is no concerning history of exertional symptoms. This would allow those with the absence of CAC to follow healthy lifestyle modifications with little or no medical therapy, whereas intense therapy is focused on a smaller population of patients with an actual higher risk of events as demonstrated by increasing atherosclerotic burden. However,

Significant Coronary Artery Disease and Artery Calcification

Significant coronary artery stenosis (> 50%) by angiography is frequently associated with the presence of CAC as assessed by EBT. The severity of angiographic coronary artery stenosis is not directly related to the total CAC. However, the absence of CAC can be extremely useful to rule out significant CAD among individuals presenting with chest pain. Our meta-analysis has shown that individuals with zero CAC are extremely unlikely to have obstructive coronary disease. ⁴¹ In our pooled analysis of 10,355 symptomatic patients who underwent coronary angiography (1941 with no CAC), the presence of CAC was highly sensitive (98%) in predicting a luminal stenosis > 50% in any coronary artery, although the specificity was low (40%). Conversely, the absence of CAC had a high negative predictive value (93%) for ruling out any clinically significant coronary

stenosis. 41 These data are consistent with direct pathological consider use of CAC measurement in asymptomatic individ-451 comparisons. 16

symptomatic patient, exclusion of measurable coronary calcium to a higher risk status on the basis of high CAC score and that may be an effective filter before undertaking invasive diagnostic subsequent patient management might be modified . However, at procedures or hospital admission." 38 Although absence of CAC is the same time, the committee did not find enough evidence for the associated with a very low likelihood of significant CAD, utility of CAC testing in further risk stratification of those approximately 2% of symptomatic individuals with significant considered at low risk as well as of those considered at high risk CAD do not have evidence of CAC. These individuals (ie, for CAD in the next 10 years. significant CAD without CAC) tend to be younger than 50 years. 44,45 As a result, one must exercise caution when evaluating younger eligibility criteria for aggressive lipid-lowering management patients for significant CAD in the absence of CAC.

to predict the absence of a significant stenosis, the ability of CAC to recent guidelines have not recommended CAC testing for those predict myocardial ischemia by myocardial perfusion imaging is with a 10-year estimated risk of < 10% (low 27 risk), 38 evidence is also encouraging although somewhat more modest. CAC scores below 100 are associated with a very low risk of cardiac ischemia, following reasons. If these guidelines are followed, most 46 which increases dramatically, especially when CAC scores are nondiabetic women who are younger than 70 years would not be 400 or higher; however, the prevalence of other clinical conditions, candidates for further risk stratification with CAC testing, such as metabolic syndrome or diabetes, may lower the threshold whereas the majority of men above 60 years will be candidates for identification by CAC of an increased likelihood of myocardial for further risk stratification. 50 Thus, a large number of women ischemia. ⁴⁷ In nine studies examining 4870 patients referred for at higher risk for CAD risk may never become candidates for perfusion stress testing (1225 with no CAC), just 6% had evidence CAC testing. At the same time, the practicality of conducting of ischemia. 41 The recent ACC/ASNC appropriateness criteria additional screening for all such women becomes an important state that a low calcium score (especially in the absence of CAC) question. The important point is that those at < 10% 10-year risk generally precludes the need for assessment by myocardial of CHD are frequently at significant longer term risk of CHD, perfusion imaging. 48

Ruling Out Acute Coronary Syndrome in the **Absence of Coronary Artery Calcification**

A pooled meta-analysis was composed of 431 patients complaining of acute chest pain with negative troponins and equivocal ECG findings based on three published studies. 41 The cohort consisted of 48% men (mean age, 51.4 years). There were only 2 of 183 patients (1.1%) without any CAC who were diagnosed with an acute coronary syndrome (ACS). Of the 248 patients with a positive CAC score, 77 (31%) were found to have an ACS. Overall, a positive CAC score had 99% sensitivity, 57% specificity, 24% positive predictive value, and 99% negative predictive value for the evaluation of ACS. The high sensitivity and negative predictive value may allow early discharge of those patients with nondiagnostic ECG findings and negative CAC score (score = 0). Long-term follow -up of this cohort demonstrates that patients without CAC at the time of the emergency visit are at very low risk of subsequent events. 49

However, the current state of the literature is certainly small and inconclusive with respect to this important clinical entity. Whereas CAC can serve as a useful marker for the exclusion of ACS in patients presenting to the emergency department, further studies in larger cohorts need to be done to establish its role in a clinical paradigm, especially with the excellent depiction not only of coronary anatomy but also of left ventricular function by contrastenhanced coronary CT angiography. 41

WHO ARE CANDIDATES FOR **CORONARY ARTERY CALCIUM TESTING?**

In 2007, the expert consensus document by the American College of Cardiology Foundation and the American Heart Association made recommendations to provide a perspective on the current state of the role of CAC testing in clinical practice. The consensus was that it may be reasonable to

als who are at intermediate risk by the FRS. 38 This conclusion was In fact, recent ACC/AHA guidelines also consider that "for the based on the possibility that such individuals might be reclassified

As far as those with 10-year risk > 20%, they already meet with optional LDL-C goals of < 70 mg/dL, and further CAC testing Whereas the absence of calcification shows exceptional ability may not lead to changes in treatment goals. However, although accumulating that such an approach may be problematic for the warranting that this issue be revisited.

> One approach would be to identify a subgroup of women in a low-risk group who are more likely to harbor significant CAC. Testing them may potentially be a cost-effective approach. Emerging evidence has strongly implicated the family history of premature CAD to be an independent risk factor strongly associated with higher burden of subclinical atherosclerosis 51-53; however, a positive family history of CAD does not factor into most global risk algorithms, such as the FRS. Nasir and coworkers 51 demonstrated that among those with premature family history of CAD (especially with sibling history), nearly one third to one guarter of self-referred patients with no or one CAD risk factor had CAC > 100. In a similar fashion, in the MESA cohort, at least 25% of individuals with family history of premature CAD had significant CAC. 52 In addition, it has been shown that among women with a family history of premature CAD along with multiple metabolic risk factors, a subgroup of women with FRS < 10% will have significant atherosclerosis. 53

> Alternatively, based on the 2003 American College of -Cardiology Bethesda Conference on atherosclerosis imaging, intermediate-risk groups could be reclassified as those at 6% to 20% risk, at least for women. 54 This strategy will identify more higher risk women who would not be placed in the intermediaterisk group, thus with lower thresholds for risk factor modification, especially regarding LDL-C control. Such persons may benefit from aggressive preventive strategies, such as statin, aspirin, and possibly blood pressure-lowering therapies, especially subsets in which the absolute risk of CAD events is increased, such as those who additionally have increased levels of CAC. Figure 27-6 provides a schematic algorithm to identify ideal candidates for CAC testing.

TRACKING PROGRESSION OF **CORONARY ARTERY CALCIUM SCORES**

A proposed use of CAC screening is to track atherosclerotic changes over time by serial measurements. There are several published studies of outcomes related to CAC progression. The first retrospective study demonstrated in 817 persons that CAC progression was greatest in those who experienced



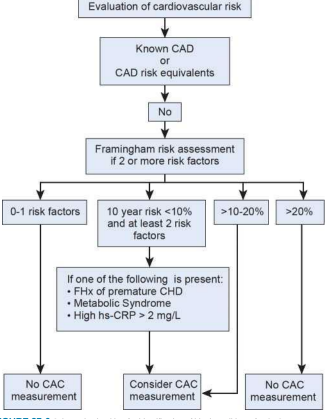


FIGURE 27-6 Schematic algorithm for identification of ideal candidates for CAC screening.

a future MI. 55 A second study measured the change in CAC in 495 asymptomatic subjects submitted to sequential CT scanning. The associated relative risk for acute MI for patients exhibiting > 15% CAC progression was elevated 17.2-fold (95% CI, 4.1 to 71.2) compared with those without CAC progression (P < 0.0001).

A large prospective study using CT to measure progression of CAC has also been reported. This study was designed as a clinical trial to evaluate the impact of aggressive lipid-lowering and antioxidant therapy on the progression of CAC in 4613 asymptomatic persons aged 50 to 70 years, with EBT scanning of the coronary arteries at baseline and again at 2 years and followup for 4.3 years. ³¹ Whereas the intervention failed to significantly affect progression of CAC, those who sustained a coronary event had a median increase in CAC score of 247 compared with only 4 in those who did not sustain a coronary event at any time during the study. Multiple logistic regression demonstrated that only age (P = 0.03), male gender (P = 0.04), LDL-C (P = 0.01), HDL-C (P = 0.04), and 2-year change in calcium score (P = 0.04) 0.0001) were significantly associated with subsequent CAD events. Increasing calcium scores were most strongly related to coronary events in this clinical study, similar to previous observational studies.

The effect of therapeutic intervention, especially statins, on the rate of progression of CAC is controversial. The earlier published retrospective and prospective cohort studies suggested a reduction in the rate of CAC progression. Callister and associates, ⁵⁶ who retrospectively studied 149 asymptomatic patients referred for sequential EBT scans at least 1 year apart, demonstrated a 45% slowing in the rate of CAC progression in those receiving statins. Budoff and coworkers, ⁵⁷ in one of the first prospective studies in this regard, demonstrated that the rate of CAC progression was 39% in those patients with dyslipidemia versus 15% in those receiving statins, representing a 61% decrease in the rate of progression with statin treatment. Similarly, Achenbach and colleagues ⁵⁸ showed that with a standard dose of 0.3 mg/day of open-label cerivastatin in 66

dyslipidemic patients, the median annual relative increase in CAC scores was 25% during the untreated period before study entry versus 9% during the treatment period (P < 0.0001). Reduction of CAC score was most prominent in those patients who achieved an LDL level < 100 mg/dL.

However, to date, these results have not been replicated in randomized controlled trials. The SALTIRE trial (Scottish Aortic Stenosis and Lipid Lowering Trial, Impact on Regression) randomized 102 patients to atorvastatin or placebo and assessed CAC progression during an average follow-up of 2 years. Despite a significant reduction in LDL and C-reactive protein levels, there was a nonsignificant increase in percent age CAC progression (26%/year with atorvastatin versus 18%/year with placebo). 59 Schmermund and coworkers 60 also failed to show reduced progression of CAC in 366 asymptomatic patients randomized to either 10 mg or 80 mg of atorv astatin during 12 months. This was despite a 20% additional reduction in LDL level in the 80-mg atorvastatin treatment group. Similarly, the BELLES (Beyond Endorsed Lipid Lowering with EBCT study, Scanning) which randomized hyperlipidemic postmenopausal women to atorvastatin 80 mg or pravastatin 40 mg, found no effect on CAC progression in both arms. Although atorvastatin reduced LDL concentration by 47% ± 20% and pravastatin reduced LDL by 25% ± 19%, there was no significant difference in CAC progression after 12 months, with an increase of 15% and 14% in CAC scores noted in the atorvastatin and pravastatin arms, respectively. 61

The St. Francis Heart Study is the only trial relating statin therapy and CAC progression to cardiac events. 62 This was a double-blind, randomized, controlled trial of atorvastatin 20 mg daily, vitamin C 1 g daily, and vitamin E (alpha tocopherol) 1000 units daily versus matching placebos in 1005 patients with coronary calcium scores at or above the 80th percentile for age and gender. Despite significant reductions in LDL and triglyceride levels during a mean treatment course of 4.3 years, there was no effect on CAC progression. However, there was a significant 42% reduction in events in treated patients who had CAC scores > 400 (8.7% versus 15.0%; P = 0.046). 62

We believe more data are needed to justify the incremental population exposure to radiation and the cost associated with a repeated CT test to assess "change" until it is better under stood what therapies may be of benefit and how clinicians should use these data in clinical practice. MESA and other population-based studies are evaluating the prognostic value of changes in CAC. Until then, consensus guidelines do not support serial measurement of CAC for the purposes of tracking the effects of therapeutic interventions. ³⁸

CORONARY ARTERY CALCIUM TESTING AND DOWNSTREAM MEDICATION INITIATION AND ADHERENCE

Adequate control of risk factors with behavioral modification and medications to control lipids, blood pressure, and other risk factors in asymptomatic patients has been the cornerstone of preventive efforts to reduce the occurrence of cardio vascular events. Among individuals with higher CAD risk, such as those with elevated CAC, these established preventive therapies are appropriate interventions. Emerging evidence also indicates that CAC may be associated with initiation of and improved adherence with cardioprotective medications. For example, Wong and coworkers, ⁶³ in a

retrospective study of 703 self-referred adults, reported that the extent of CAC is independently associated with beginning of lipid-lowering medications and aspirin as well as with initiation of healthy lifestyle changes, including losing weight and decreasing dietary fat. Taylor and colleagues, 64 who compared statin and aspirin use in 1640 young men, reported that at 6-year follow-up after a CAC test, statin use was threefold more likely among those with any CAC compared with CAC = 0 (48% versus)15%; P < 0.001) and aspirin use was nearly twice as likely (53%) versus 32%; P < 0.01). Our group has also previously shown that those with elevated CAC scores are almost three times more likely to initiate aspirin therapy compared to subjects with absent CAC scores. 65 Overall, aspirin initiation was lowest (29%) among those with CAC = 0 and gradually increased with higher CACscores (1 to 99, 55%; 100 to 399, 61%; > 400, 63%; P < 0.001 for trend). In a multiethnic cohort, Nasir and colleagues 66 demonstrated that after demographics, CAD risk factors, and socioeconomic factors are taken into account, a high CAC score (> 400) is associated with a 32% to 55% higher likelihood of initiation of these cardioprotective medications in a mean followup of 1.6 years.

Although there appears to be a consistent relationship of elevated CAC scores with initiation of preventive therapies, data on whether atherosclerosis imaging improves medication adherence are conflicting. Our group reported in a study of 505 asymptomatic individuals that continuation of lipid-lowering medication was lowest (44%) among those with a CAC score in the first quartile (0-30), whereas 91% of individuals with a baseline CAC score in the fourth quartile (> 526) adhered to lipid-lowering medication. 67 In multivariable analysis, after adjustment for cardiovascular risk factors, age, and gender, higher baseline CAC scores were strongly associated with adherence to statin therapy. However, no significant differences were noted in continuation of either lipid-lowering medication (87% versus 79%; *P* = 0.06) or aspirin (79% versus 83%; *P* = 0.36) according to presence or absence of CAC. Nasir and colleagues 66 showed that the risk ratios for medication continuation with elevated CAC scores were 1.10 (95% CI, 1.01-1.20) for lipidlowering medication, 1.05 (1.02-1.08) for blood pressurelowering medication, and 1.14 (1.04-1.25) for aspirin initiation. Moreover, in the only published randomized clinical trial assessing the effects of CAC scanning on estimated risk of CAD after 1 year determined by changes in FRS, 68 there was no difference in mean absolute risk change in 10-year FRS (+ 0.30 versus + 0.36; P = 0.81) comparing overall groups who received CAC score results with those who did not; however, those who received intensive case management of risk factors versus usual who had a significantly better outcome (change in FRS of - 0.06 versus + 0.74; P = 0.003). In this study, the prevalence of CAC was fairly low (15%), with generally low CAC scores in those with CAC; thus, it is possible that power was limited to impact on change in risk among most of the participants who did not have CAC.

In general, we think that CAC assessment not only may have a role in the diagnosis of coronary atherosclerosis but also may improve the likelihood of individuals at highest risk of coronary events to be on appropriate preventive medication. Emerging evidence indicates that the rate of use of these medications in the group that would most likely benefit from lipid-lowering therapy (those patients with increased levels of subclinical atherosclerosis) is substantially higher. The results of these studies in general suggest that CAC found on cardiac CT may add much needed motivation to asymptomatic patients recommended for lifestyle modification and drug therapy; however, properly designed randomized trials remain necessary to evaluate the true role of CAC testing in improving clinical outcomes that may be a result of any such risk factor modification efforts.

CORONARY ARTERY CALCIUM TESTING AND **DOWNSTREAM TESTING AND COSTS**

From a societal standpoint, apart from the ability of CAC testing to predict future CAD outcomes, a key point to be demonstrated is that atherosclerosis imaging for adults will not lead to a cascade of costly downstream testing. This issue becomes even more relevant in the current economic climate, and there are active governmental efforts to curtail health care expenditures. 69 Shaw and coworkers 70 showed that the majority of individuals who had either no CAC or minimal CAC scores of 1 to 10 had very few downstream additional cardiac tests, whereas the majority of testing was performed in those with advanced CAC. Noninvasive testing was infrequent and medical costs were low 27 among subjects with low CAC scores, both rising progressively with increasing CAC scores (P < 0.001), particularly in the 31 (2.2% of subjects) who had CAC scores > 1000. Similarly, invasive coronary angiography rose progressively with increasing scores (P < 0.001) but occurred exclusively among subjects first undergoing non-invasive testing and overall was performed in only 19.4% of subjects with CAC scores > 1000. It was clearly shown that invasive procedures were not performed immediately after CAC testing, and they were performed in a stepwise manner preceded by functional imaging with either exercise stress testing or stress myocardial perfusion imaging. 70

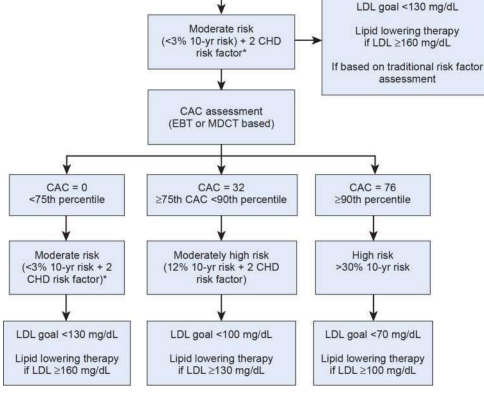
On the basis of our assumptions derived from the MESA study, among every 100 individuals screened for CAC, only 8 would have CAC scores > 400, with the majority of these possibly undergoing stress echocardiography or myocar dial perfusion testing (5 to 7 individuals) and nearly half undergoing invasive coronary angiography (approximately 4 individuals). In comparison, only 14 individuals of the 78 of 100 screened with CAC scores of 0 to 10 would undergo some sort of stress imaging, and only 1 will proceed with invasive coronary angiography in 6 years of follow-up. For the majority of individuals with no or low CAC (78% of those screened with CAC), the median costs were minimal (\$25 to \$35), mostly incurred by ECG testing, which is often part of the initial assessment of individuals with hypertension (a feature observed in nearly 60% of this low-risk group). 69

Although the emerging data are encouraging, we hope that the various stakeholders in determining health care resource allocation will quickly move in the direction of addressing whether selective use of atherosclerosis imaging should play any role in halting the epidemic of atherosclerotic vascular disease by better refining which middle-aged and older adults are truly at relatively high risk versus very low risk for a CVD event during the next 5 to 10 years. There is an urgent need of a randomized trial that compares the current traditional risk factors-based approach with one supplemented by sub-clinical atherosclerotic screening to determine whether this approach can save lives in a manner that is at least moderately cost-effective.

USING INFORMATION FROM CORONARY ARTERY CALCIUM TESTING TO MODIFY TREATMENT

Incorporation of CAC into preventive strategies applied in the context of conventional clinical profiles can potentially refine risk definition or understanding of the consequences of risk factors into a more complete or comprehensive assessment in any given person. At the moment, we have proposed that the CAC scoring can be used to modify the number of points assigned to chronological age in determining global risk





Global CHD risk assessment

FIGURE 27-7 Incorporation of the CAC score with global risk assessment for a 40-year-old asymptomatic man, past but not current smoker, no diabetes, with total cholesterol of 205 mg/dL and HDL of 41 mg/dL, and mild untreated hypertension [systolic blood pressure of 150 mm Hq].

assessment, such as in the Framingham risk or European risk score estimates, for more accurate prediction of 10-year -cardiovascular risk. ⁷¹ The use of CAC scoring in combination with conventional risk factor assessment to define a "modified " FRS allows the inclusion of subclinical disease definition into the context of modifiable risk factors.

Merely finding subclinical disease does not imply which treatable factors require modification. Merely establishing an elevated blood cholesterol level or smoking status does not imply that these factors have contributed to the actual development of disease because genetics and other unidentified factors will also influence the presence of disease in any given individual. In the FRS, age can be largely considered a proxy for the atherosclerotic burden, which increases progressively with age but may vary greatly between individuals. The employment of CAC scores to identify the "heart age" allows accounting for the variability in atherosclerotic burden at a given age and can be easily incorporated into FRS as well as the NCEP guidelines.

An example of the incorporation of the CAC score with estimation of global risk score is shown in Figure 27-7. For example, in a person determined to be at low risk by global risk assessment, consistent with high event rates associated with higher CAC burden, it seems appropriate to initiate drug therapy at lower thresholds of LDL-C (eg, 130 mg/dL instead of 160

mg/dL) in those with CAC scores in the > 75th percentile group and at even lower cut points (eg, 100 mg/dL) when CAC is > 90th percentile for age or gender in the presence of multiple risk factors. This is especially true among individuals with a strong family history of premature CAD as they tend to have higher CAD event rates even at lower levels of predicted risk. In these persons, especially with CAC scores > 400, further evaluation by functional testing, such as myocardial perfusion imaging, may be considered.

Use of cardiac CT screening, even at as high a cost as \$500 per study, will still be about half the cost of a brand name statin therapy for a year. In recent years, many cities and academic centers charge \$100 or less for a coronary calcium scan. Identification of those with no or very mild CAC as those in whom a more conservative LDL goal may be acceptable has been previously suggested to provide a cost-effective use of EBT/CT technology in clinical practice. ⁷²

CONTRAST-ENHANCED COMPUTED TOMOGRAPHY: ASSESSMENT OF NATIVE CORONARY ARTERIES **CORONARY**

^{*} If only one risk factor, then LDL goals <160 mg/dL in presence of 10-year risk <10%

components). 73 As the reliability of MDCT in the detection and grading of coronary artery stenosis has been established, the times, reduced breath-hold duration, smaller injections of challenge now is to define the ability of this diagnostic tool to intravenous contrast material, and decreased motion-related distinguish the coronary artery plaque prone to rupture with artifacts, resulting in lower radiation exposure and improved findings suggestive of instability. In the following section, we diagnostic accuracy. describe briefly the image acquisition and analysis sessions and mainly elucidate our current understanding, explore the Image Interpretation relationship of plaque subtypes with current and future cardiovascular risk, and assess the potential role of CTA in primary preventive strategies.

Image Acquisition

MDCT scanners produce images by rotating an x-ray tube around can be limited by overlapping structures adjacent to the artery of a circular gantry through which the patient advances on a moving interest. Curved multiplanar reformations are reconstructed on a table. Pitch is the speed of the table relative to the speed of the plane to fit a curve and allow display of the entire vessel in a single gantry rotation, which allows each cross sectional level of the heart image. Three dimensional volume-rendered images are useful to be imaged during more than one cardiac cycle. The number of for selection of images with the least motion artifact and for image slices acquired during each gantry rotation (4 to 320) assessment of the relationships among different anatomical determines the overall duration of the MDCT scan. Developments structures. Most assessments of plaque, however, have been in MDCT technology have led to the rapid advancement from 4- qualitative in nature, and there is yet to be any reliable software slice MDCT machines in 1998 to 64-multislice scanners available for that can reproducibly quantify the amounts of soft or mixed clinical use in 2004. ²⁴ The current 64-slice MDCT systems are plaque burden and that is not cumbersome to use. capable of simultaneously acquiring 64 sections of the heart with the fastest gantry speeds of 330 msec per rotation. The increased Association of CTA-Detected Plaque Subtypes numbers of detectors have greater craniocaudal coverage per with Traditional Risk Factors rotation, which results in faster scan times and allows the entire cardiac anatomy to be imaged in less than 10 seconds.

Cardiac CT is performed with ECG gating in a prospective or retrospective mode. ECG gating synchronizes image acquisition with the cardiac cycle. The optimal phase or interval for image analysis is the period during which the heart is the least mobile (usually end diastole) and the least degraded by motion artifact. Prospective ECG gating entails scan initiation at a defined interval after the R wave, continues for a pre-specified duration, and then stops until the same optimal period is reached in the subsequent right coronary artery. Few studies have assessed the relationship cardiac cycle, at which time scanning resumes.

Retrospective ECG gating employs continuous acquisition of images throughout the cardiac cycle. The images from multiple consecutive heartbeats are then reconstructed at various percentages of the RR interval (eg, from 0% to 90% of the RR cycle at 10% intervals). With retrospective gating, several thousand images can be acquired during a single cardiac study, allowing the interpreting physician to select the images with the least amount of presence of any plaque as well as for an increased burden of motion-related distortion before final image reconstruction. Gating is the most advantageous at slow heart rates (less than 60 beats/min), when the RR interval is > 1000 msec and the fastest imaging protocols are used.

Cardiac motion is minimized with the oral or intravenous administration of beta blockers before scanning, thereby reducing the heart rate and prolonging the time during the cardiac cycle at which coronary artery velocity is low. For those individuals without contraindication to beta blockade, these drugs are the medication of choice because they not only decrease the heart rate through the reduction of sympa thetic tone but also may reduce the postulated that underlying coronary plaque differences may in number of premature atrial or ventricular beats, which adversely affect the overall quality of the images. Another crucial element for high-quality coronary images to be obtained is maximal dilation of the coronary vessels with nitroglycerin through the use of sub lingual tablets or spray. Respiratory motion is excluded by performance of the scan during a breath-hold.

Coronary CTA requires the intravenous administration of an compared with iodinated contrast medium (which is contraindicated in persons with renal impairment). Approximately 50 to 100 mL of contrast medium is necessary for adequate

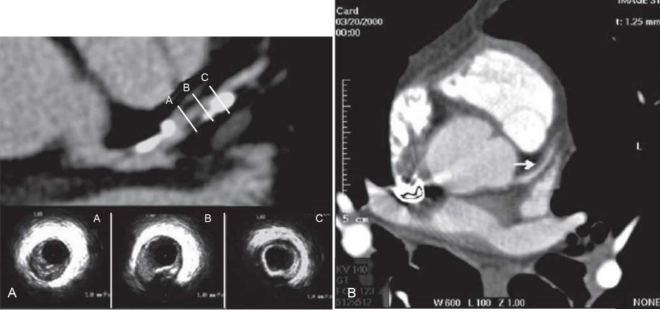
coronary artery enhancement. The accurate timing of image 455 acquisition relative to the injection of contrast material is a major determinant of overall image quality. A test bolus or bolus tracking technique is used to optimize this timing by determining the amount of time necessary to peak enhance ment in the aorta.

Advancements in MDCT technology have led to shorter scan

The coronary vasculature on cardiac CTA is evaluated through 27 axial images, multiplanar (coronal, sagittal, or oblique) reforms, and three-dimensional data sets constructed from specific phases during the cardiac cycle. Maximum intensity projection images allow the evaluation of longer segments of the coronary vessels but

There has been increasing interest in the detection and quantification of plaque subtypes by CTA, including calcified, mixed, and noncalcified plaque. Figure 27-8 shows examples of CTA-assessed calcified and noncalcified plaque, including correlation with intravascular ultrasound (Fig. 27-8A). Figure 27-9 shows a volume-rendered image demonstrating severe coronary atherosclerosis affecting the distal left main, proximal left anterior descending, and first diagonal and the proximal of traditional risk factors with coronary plaque subtypes determined on CTA. ⁷⁴⁻⁸⁰ Rivera and associates ⁷⁴ reported on the relationship of traditional cardiovascular risk factors with the presence and burden of plaque subtypes in more than 1000 asymptomatic Korean individuals who underwent CTA. In agreement with what has been previously shown with CAC, age and male gender were overall the strongest predictors for the calcified, mixed, or noncalcified plaque. The study found that specific plaque types are more strongly associated with certain risk factors. For example, age and hypertension were the strongest predictors of mixed plaque burden, whereas smoking was strongly associated with the burden of noncalcified plaque; LDL-C related only to the presence and burden of mixed plaque.

It is also well known that individuals with diabetes have a significantly higher risk for the development of coronary events. For this reason, there is increased interest in the non-invasive assessment of coronary plaque in this population of patients. It is part explain the increased risk of CVD in this vulnerable group. Our group assessed 416 symptomatic patients (64% men; mean age, 61 ± 13 years), with 61 (15%) reporting type 2 diabetes, who underwent CTA. Enrolled patients had an intermediate pretest probability of obstructive CAD. 77 Patients with diabetes had a higher number of coronary segments with mixed plaques



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FIGURE 27-8 A, Correlation of CTA of the coronary arteries with intravascular ultrasound illustrates the ability of MDCT to demonstrate calcified and noncalcified coronary plaques. B, Noncalcified, soft, lipid-rich plaque in left anterior descending artery (arrow) (Somatom Sensation 4, 120 mL Imeron 400). The plaque was confirmed by intravascular ultrasound. (A from Becker CR, Ohnesorge BM, Schoepf UJ, Reiser MF: Current development of cardiac imaging with multidetector-row CT. Eur J Radiol 36:97, 2000. B from Kopp AF, Kuttner A, Trabold T, et al: Multislice CT in cardiac and coronary angiography. Br J Radiol 77:S87, 2004. Reproduced with permission.)



FIGURE 27-9 Volume-rendered image from 16-slice Toshiba Aquilion multidetector scanner showing distal right coronary artery and veins. Severe coronary atherosclerosis, with significant obstructions affecting the distal left main, proximal left anterior descending, and first diagonal and the proximal right coronary artery. (Courtesy of Toshiba, Inc.)

nondiabetic patients (1.67 \pm 2.01 versus 1.23 \pm 1.61; P = 0.05), whereas no such differences were observed for noncalcified and calcified components. Even after traditional risk factors were taken into account, patients with diabetes were more likely to have an elevated burden (three or more segments) of mixed coronary plaque (odds ratio, 2.34; 95% CI, 1.14 4.83). On the other hand, no significant burden association was observed with noncalcified (0.38; 95% CI, 0.08-1.71) or calcified (0.98; 95% CI, 0.50-1.95) plaques.

Similar findings have also been reported by Rivera and

associates, 78 who studied 217 asymptomatic Korean outpatients with type 2 diabetes who had no prior history of CAD. A total of 138 individuals (64%) had occult CAD (any plaque) on CTA. Similar to prior reports, most subjects (62 of 138; 45%) had a combination of noncalcified, calcified, and mixed coronary plaques, whereas exclusively noncalcified, calcified, and mixed plaques were seen in 9% (13 of 138), 20% (27 of 138), and 26% (36 of 138), respectively. ⁷⁸ Rivera and associates ⁷⁹ also investigated the association of coronary plaque subtypes with the level of glycosylated hemoglobin in a group of 906 asymptomatic individuals without diabetes. Unadjusted analyzes demonstrated a positive association between increasing levels of hemoglobin A1c and the number of coronary segments with mixed plaques (P <0.0001). After traditional risk factors were taken into account, those with hemoglobin A1c in the highest tertile (5.9% to 7.0%) versus the lowest tertile (< 5.4%) had a relative risk of 4.6 (95% CI, 1.3-16.5) of having two or more segments with mixed plaque. No relationship was seen with calcified or noncalcified plaques.

Another key area of interest that relates to coronary plaque morphology is the difference in plaque composition between men and women. This could potentially help determine the natural history of atherosclerosis in the two genders as well as pinpoint the possible difference in the prevalence of plaque subtypes. These findings could be relevant, taking into consideration that women at almost all age groups are at a lower risk of experiencing cardiovascular events. Few reports based on intravascular ultrasound studies have identified sex differences in the morphology of coronary artery plaques, suggesting that coronary plaques in women contain relatively large amounts of cellular tissue and relatively little dense fibrous tissue and calcium. However, these

evaluations are limited by the invasive nature of the procedure as enhanced MDCT. These findings are consistent with prior well as by limited assessment of few coronary segments and thus invasive reports that more plaques with tiny calcific deposits are may not reflect differences in the overall plaque burden. Nasir and noted with culprit lesions in ACS patients.³³ coworkers 80 demonstrated that among symptomatic individuals undergoing CTA, women presented with a significantly lower mean number of segments containing calcified plaques (1.43 ± 2.04 Association of CTA-Detected Plaque Subtypes versus 2.25 \pm 2.30; P = 0.004) as well as mixed plaques (1.67 \pm 1.23 with Cardiovascular Disease Outcomes versus 2.25 ± 2.30 ; P = 0.05), whereas no such relationship was seen with noncalcified plaques (0.72 \pm 1.01 versus 0.86 \pm 1.06; P = 0.21). On the other hand, the relative proportion of overall plaque burden was more likely to be noncalcified (40% versus 28%) and less likely to be mixed (22% versus 28%) as well as calcified (38% versus 43%). In other words, despite similar overall noncalcified plaque burden, women were found to have a higher proportion of noncalcified to total plaque (ie, larger relative burden of noncalcified plaque). The potential mechanisms of the higher proportion of noncalcified plaque in women are not entirely clear, and further research is needed into whether endogenous hormonal factors, such as estrogen, may play a role.

Association of CTA-Detected Plaque Subtypes with Myocardial **Perfusion Defects**

Detection of myocardial perfusion abnormalities in a non-invasive manner by modalities such as single-photon emission computed tomography (SPECT) is a method for identification of high-risk patients and stratification according to cardiovascular prognosis. In one study, Lin and colleagues 81 evaluated 163 consecutive low- to intermediate-risk symptomatic patients without known CAD who underwent both stress SPECT and MDCT. Overall, 33 (17%) had a summed stress score of > 8 (abnormal myocardial perfusion). Among individuals with a summed stress score > 8 versus < 8, a significant difference only in the number of mixed plaques was reported (2.10 \pm 2.50 versus 1.16 \pm 1.69; P = 0.01), whereas no differ ences were seen in noncalcified (1.03 \pm 1.67 versus 0.97 \pm 1.44; P =0.83) and calcified (2.10 \pm 2.50 versus 1.56 \pm 1.59; P = 0.54) plaques. In multivariate analyses, adjusting for traditional risk factors, mixed plaque burden remained associated with summed stress score > 8 (odds ratio, 1.28; 95% CI, 1.05-1.56). Although the study findings imply a potential role of assessment of mixed plaque burden, studies are needed to confirm these findings in large heterogeneous populations.

Association of CTA-Detected Plaque Subtypes with Features of Plaque Vulnerability on **Intravascular Ultrasound Examination**

Autopsy studies have demonstrated that individuals suffering from sudden cardiac death have multiple plaques containing a large amount of necrotic core with an overlying thin-cap fibroatheroma, which are traits believed to be linked to plaque on the extremes (non calcified and calcified), but the eventual rupture. 14 Although intravascular ultrasound is well equipped to prognostic importance of mixed plaque may eventually be seen assess plaque subtypes as well as thin-cap fibroatheroma, whether in future long-term studies. certain plaque types now detected on coronary MDCT provide clues to these vulnerability features is not well studied. In a landmark study, Pundziute and coworkers 82 assessed the RADIATION DOSE prevalence of plaque subtypes on MDCT and the relationship with thin-cap fibroatheroma based on intravas cular ultrasound in 25 patients with ACS. In these vulnerable patients, 32% of plaques were noncalcified on MDCT and 59% were mixed. More interestingly, intravascular ultra sound-derived thin-cap fibroatheromas were most frequent

observed in lesions classified as mixed (68%) compared with 457 noncalcified (19%) and calcified (13%; P = 0.001) on contrast-

On non-contrast-enhanced cardiac CT, the identification and quantification of CAC provide prognostic information incremental to conventional risk factors in predicting adverse events | cardiac events. There is a significant lack of knowledge about 27 the "prognostic" significance of noncalcified versus calcified plaque detectable with CTA on top of overall plague burden assessment, for which the CAC score is an excellent surrogate. To date, to the best of our knowledge, only one study has attempted to assess the relationship of coronary plaque subtypes detected by MDCT and future events. Pundziute and coworkers 83 observed a total of 100 symptomatic patients who underwent coronary MDCT for the occurrence of cardiac death, nonfatal MI, unstable angina requiring hospitalization, and revascularization. During a mean follow-up of 16 months, 33 events occurred in 26 patients. Among all plaque subtypes, only mixed plaque burden was associated with adverse events (hazard ratio, 1.6; 95% CI, 1.6-2.0).

In recent years, the development of contrast-enhanced coronary CTA has enabled the identification of different plaque subtypes (exclusively calcified or mixed plaques) and has generated great enthusiasm, given the potential of identifying vulnerable plaques." The emerging literature provides insights into the relationship between plaque char acterization with MDCT and varying clinical scenarios, suggesting that plaque composition could have incremental value and potentially be included in overall risk stratification strategies. Currently, it is not entirely clear whether exclusively calcified or noncalcified plaque subtypes predispose to higher cardiovascular risk; however, data from the limited studies in this regard suggest that mixed plaque burden is more likely to be associated with highrisk groups, such as patients with diabetes mellitus and those with elevated inflammatory biomarkers. In addition, mixed plaque has been shown to be strongly associated with myocardial perfusion defects and features of plaque instability such as thincap fibroatheroma. Recently, CT-assessed characteristics such as positive vessel remodeling and low-attenuation plaques have also been shown to be associated with subsequent development of ACS, 84

Although one small study has suggested mixed plaque burden to predict future outcomes, it is not entirely clear if there is additional prognostic value above and beyond the CAC score, an established marker of future CAD events, especially in the absence of detectable CAC. 85 We believe that large long-term studies are needed to clearly elucidate the role of these plague subtypes in the overall risk assessment strategies. Until then, we believe the value of plaque assessment by CTA appears to be less

Increased awareness of radiation in the medical community and general population has raised warnings about unnecessary sary testing, especially in the context of CTA, for which the mean doses are rather high compared with some other forms

458 of cardiac testing. Radiation doses with coronary calcium scores are sufficiently low (approximately 1 mSv) with either EBT or MDCT, which is far below background annual radiation exposures. ²⁴ CTA doses for retrospective triggering are in the range of 10 to 18 mSv (which is similar to cardiac nuclear doses). Since the increased awareness and concern about radiation, several radiation dose reduction techniques have been introduced. These include dose modulation (lower radiation dose 30% to 48%), reduction of kilovoltage to 100 kVp for thinner patients (lower radiation dose by 40%), E limitation of the top and bottom of the scan field (lower radiation dose by 20%), ⁸⁶ and prospective triggering (lower radiation dose by 70%). ⁸⁷ Collectively, the dose reduction will be approximately 80% to 90%.

27 One study, without use of any other dose reduction technique except for prospective imaging, reported that the mean radiation dose to the patient was 77% lower for prospective gating (4.2 mSv) than for retrospective gating (18.1 mSv) (*P* < 0.01), without compromising image quality or diagnostic accuracy. ⁸⁸ Another study similarly reported that use of prospective triggering reduced radiation exposure by 80% without compromising image quality compared with traditional retrospective acquisition. ⁸⁹

GUIDELINES FOR CORONARY ARTERY CALCIUM TESTING AND COMPUTED TOMOGRAPHIC ANGIOGRAPHY IN ASYMPTOMATIC INDIVIDUALS

At the same time, the current guidelines do not recommend the use of CAC scanning in subjects at high risk (e..g, > 20% risk of a CAD event in 10 years) or with preexisting CAD or diabetes, for which aggressive guidelines already exist, so results of CAC scanning should not influence therapeutic decisions, or in those at low risk (eg, population screening in those with < 10% risk of a CAD event in 10 years). Moreover, serial imaging for assessment of progression of coronary calcification is not recommended, nor are there any indications for performing CTA for risk assessment in asymptomatic individuals or for monitoring disease progression in any group of patients. $^{24.38}$

CONCLUSION

Global risk assessment seems to underestimate the CAC burden as well as the CVD risk, especially in selected young men and postmenopausal women with multiple risk factors. CAC is an independent predictor of CAD events and provides prognostic information above and beyond the FRS. Assessment of coronary artery calcification with non-contrast-enhanced CT appears to be most predictive among individuals with intermediate risk. Although current recommendations do not support its use as a screening tool in low-risk patients, emerging data suggest that many of these individuals with a family history of premature CAD may potentially benefit from this testing.

At the other end of the spectrum, those with high pretest probability are essentially at CAD-equivalent risk regardless of calcium score, and treatment of risk factors rather than screening would be more appropriate. Integration of the information generated from the FRS with CAC, especially by incorporation of age as another criterion in risk assessment, appears to be an effective system for assessment of actual cardiac risk to optimally target and follow the effect of preventive measures. In addition to providing information on risk stratification, CAC testing may improve adherence to preventive medication use in appropriate high-risk individuals . To date, there does not seem to be evidence for a substantial increase in downstream testing with CAC testing in asymptomatic individuals. There is an important need, however, for randomized clinical trials to better document the clinical utility of both CAC scanning and CTA with respect to clinical outcomes.

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CHAPTER 28

Use of Cardiac Magnetic Resonance Imaging and Positron Emission Tomography in Assessment of Cardiovascular Disease Risk and Atherosclerosis Progression

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KEY POINTS

- Atherosclerosis is a chronic immunoinflammatory disorder of the vascular, metabolic, and immune systems.
- Detection of subclinical atherosclerotic disease will enable initiation of earlier treatment and prevent the progression of atherosclerotic vascular disease and plaque disruption.
- MRI can evaluate various aspects of cardiovascular disease, including plaque morphology, vascular physiology, and myocardial function.
- High-resolution multicontrast MRI is used to characterize the specific tissue components of complex plaques, which aids in the detection of vulnerable plaque.
- Because of high reproducibility, MRI is valuable in assessment of plaque progression and regression.
- Molecular imaging elucidates the biology of various cellular and molecular targets in atherosclerosis before the development of gross phenotypic changes.
- MRI is an ideal molecular imaging modality because of its high spatial and temporal resolutions, noninvasive nature, lack of ionizing radiation, and excellent reproducibility.

- PET can assess inflammatory activity within atherosclerotic plaque because of avid FDG uptake and accumulation within the macrophages in inflamed plaque.
- PET can simultaneously evaluate cardiac perfusion and metabolism and hence plays a vital role in the evaluation of selected patients with cardiovascular risk factors or disease.

Atherosclerosis is a diffuse, chronic -immunoinflammatory disease of the vascular, metabolic, and immune systems characterized by deposition of lipid and fibrous products in the arterial wall. It is the result of complex biological processes, resulting in various local and systemic manifestations after a long asymptomatic period. Genetic predisposition is part of the variability in the development of atherosclerotic disease, plaque destabilization, and subsequent thrombosis.

Atherosclerosis is the single most important contributor to the burden of cardiovascular disease. ¹ In spite of advances in prevention, risk assessment, and treatment, cardiovascular disease is the major global cause of morbidity and mortality. An esti mated 80 million American adults have car diovascular disease, which accounted for 35% of deaths and an estimated cost of \$475.3 billion dollars in 2009. ² The large population at risk for coronary events and frequent first presentation with a significant cardiovascular event or sudden death makes it seem reasonable to detect subclinical disease for initiation of earlier treatment to prevent the progression of atherosclerotic vascular disease and plaque disruption.

Conventionally, imaging has focused on detection and grading of stenotic lesions and perfusion abnormalities in organs distal to the stenosis. However, major acute events, such as myocardial infarction (MI) and stroke, are generally produced not by plaques with high-grade stenosis but by disruption of plaques that did not produce significant stenosis or perfusion abnormalities

in stress tests, the so-called vulnerable plaque. In addition, recent evidence suggests that interventional treatment of coronary stenosis does not significantly improve MI and death rates in many persons with stable angina compared with a strategy of aggressive medical and lifestyle changes. ³

Hence, there is a need for new diagnostic tests that will be able to detect, quantify, and characterize the plaque, including its biological activity and stability, ideally at an earlier subclinical stage in the disease process so that high-risk patients can be identified and treatment initiated earlier in the long asymptomatic phase to modify the disease progression and to reduce the risk of a major event. Because atherosclerosis is a diffuse disease, detection in one vascular territory implies involvement of other vascular territories, and treatment has to be both local and global. ⁴ Novel therapeutic strategies include targeted transport vehicles allowing drug delivery to specific cells or cell structures.

IMAGING OF ATHEROSCLEROSIS

The purposes of imaging in atherosclerosis are understanding of the natural history and pathobiology of atherosclerosis, diagnosis of subclinical disease and risk stratification, evaluation of plaque burden (location, size, chemical composition, biological activity), and identification of vulnerable plaque. Serial imaging is performed to assess plaque

The ideal imaging technique should be sufficiently sensitive and specific for detection of atherosclerosis, inexpensive, reproducible, easy to perform, widely available, tolerated by patients, and noninvasive or minimally invasive. It should have no or minimal radiation, provide immediate results, quantify plaque components, be feasible in all vascular beds, add predictive value over measurement of established risk factors, and correlate highly with risks of subsequent major events. 5

The various available imaging options include invasive techniques, such as coronary angiography, intravascular ultrasound, palpography, and optical coherence tomography, and noninvasive techniques, such as high-resolution ultra sound, computed tomography (CT), single-photon emission computed tomography (SPECT), positron emission tomography (PET), and magnetic resonance imaging (MRI). Each has its advantages and disadvantages.

MAGNETIC RESONANCE IMAGING

MRI uses the magnetic characteristics of the most abundant proton in the human body, hydrogen, to generate images. When they are placed in a powerful magnet, these protons align with the magnetic field; but on application of radio frequency pulses, this alignment can be altered, causing the hydrogen protons to produce a varying magnetic field that can be detected by the scanner. The signal can be manipulated by multiple, additional magnetic fields for reconstruction of an image of the tissues of interest. The combination of radiofrequency and gradient waveforms used to obtain an image is called a pulse sequence. The appearance and signal intensity of any tissue on MRI depends on the type of pulse sequence and imaging parameters.

Rapid advances in the last 10 years have resulted in the emergence of MRI as an important tool in the evaluation of car diovascular disease. This is attributed to improved hardware, particularly the high-field strength magnets and high-sensitivity coils, and novel contrast agents, including those labeled with molecular targets. MRI is noninvasive and has no ionizing radiation; it has high inherent soft tissue contrast, spatial temporal resolution, large field of view, and multiplanar imaging capabilities. Intravenous chelated gadolinium contrast material is useful for further characterization, but it is best avoided in patients with severe renal dysfunction because of the association with nephrogenic systemic fibrosis, a debilitating fibrosing condition believed to be secondary to an immune reaction to gadolinium particles deposited in the layers of skin, most probably due to high serum concentrations in patients with impaired renal function.

MRI Techniques in Cardiovascular Diseases

MRI plays an important role in the evaluation of various aspects of cardiovascular disease (Table 28-1).

High-Resolution Multicontrast Imaging

High-resolution MRI is used to characterize the atherosclerotic plaque. The normal arterial wall has three layers, namely, the tunica intima, media, and adventitia, which cannot be separately distinguished on MRI. However, plaques are composed of variable quantities of cells, connective tissue, lipids, calcification, and debris, which can be evaluated by MRI (Table 28-2, Fig. 28-1). 6,7

> TABLE 28—1 Role of MRI in Evaluation of Various Aspects of Cardiovascular Disease

Technique Assessment

MR angiography Whole-body MRA	Stenosis, dilation Screening
•	Plaque detection, quantification,
High-resolution MRI	characterization
Vascular physiology	Compliance, aortic stiffness with pulse wave velocity, flow-mediated vasodilation
Functional data	Wall shear stress, neovascularization density
Molecular imaging	Detection of cellular and molecular targets
	Assessment of vulnerable plaque
Myocardial function	
	Volumes, mass, function Regional function
Stress perfusion study	Ischemia
Delayed enhancement	Viability
Coronary MRA	Coronary luminal assessment
Peripheral arterial MRI	MRA luminal assessment High-resolution MRI for plaque characterization
	Phosphocreatine kinetics
Intravascular MRI	Plaque assessment and characterization Guidance for coronary intervention

Signals on MRI depend on proton density and T1 (longitudinal) and T2 (transverse) relaxation times of protons in water and fat. Differences in relaxation times of different tissues produce the contrast in MRI images, which can be manipulated by changing scan parameters or with contrast agents that modify relaxation times. MRI has high inherent soft tissue contrast that is ideal for characterization and differentiation of plaque components on the basis of biophysical and biochemical parameters, such as chemical composition, concentration, water content, physical state, molecular motion, and diffusion.

High-resolution MRI with multicontrast black blood techniques and dynamic contrast-enhanced angiography is used to characterize the specific tissue components of complex plaques, which aids in the detection of vulnerable plaque. High field magnets (1.5T or above) and phased array surface coils are used. Electrocardiography-gated axial images of the thoracic aorta are acquired in expiratory breath-hold. Multi contrast techniques use a combination of black and bright blood techniques that usually include T1 weighting, T2 weighting, proton density weighting, and three-dimensional time of flight, followed by postcontrast T1 weighting and time of flight.

Black blood images are acquired by the double inversion recovery fast spin-echo technique, 8 which eliminates signal from flowing blood, resulting in high contrast between the dark blood and arterial wall with signal, enabling precise identification of the lumen-wall interface that is critical for assessment of plaque composition. T1, T2, and proton density weighting are achieved by using different relaxation time (TR) and echo time (TE). Magnetization transfer and diffusion imaging are used in some protocols. 9 Combinations of these different weightings help distinguish all the major plaque components, such as the lipidrich necrotic core, fibrous cap, loose matrix, calcification, and intraplaque hemorrhage, all of which have different signal characteristics (Table 28-3). 6,10 The MRI findings have been validated in histological studies and are highly reproducible both qualitatively and quantitatively, thus making MRI useful for serial evaluation of plaque progression or regression.

Stage **Histologic Appearance** MRI Stage MRI Appearance Not grossly apparent; isolated macrophages with oxidized lipid droplets (foam Normal wall thickness; thin plaque, no calcification, <10% stenosis Ш Fatty streak with multiple foam layers Ш Ш Diffuse intimal thickening; small eccentric plaque, no calcification Preatheroma: raised fatty streak; multiple, small extracellular lipid cores; foam cells with lipid droplets; increasing smooth muscle cells IV IV-V Plaque with lipid or necrotic core, covered by fibrous cap, possible Atheroma: single, massive extracellular lipid pool, covered by proteoglycansmall calcifications rich laver infiltrated with foam cells and smooth muscle cells with and without lipid droplet inclusion V Fibroatheroma: type IV, but with a surrounding cap rich in fibrosis, possible small calcifications VI Complex plaque with possible surface defect, hemorrhage, or Complex plaque with possible surface defect, hemorrhage, or thrombus thrombus

TABLE 28—2 Conventional and Modified American Heart Association Classification of Atherosclerotic Plaques for Histologic Diagnosis and MRI

Modified from Cai JM, Hatsukami T, Ferguson MS, et al: Classification of human atherosclerotic lesions with in vivo multicontrast magnetic resonance imaging. Circulation 106:1368, 2002.

VII

VIII

Calcified plaque

Fibrous tissue, no lipid core, possible small calcifications

VII

VIII

Calcified plaque

Fibrotic plague without lipid core; fibrous tissue, no lipid core

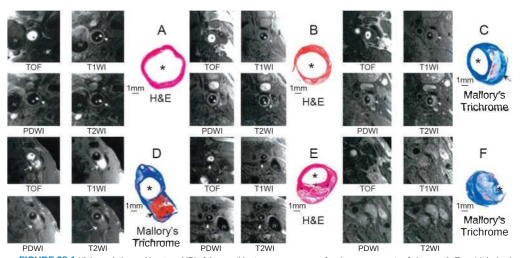


FIGURE 28-1 High-resolution multicontrast MRI of the carotid artery: appearances of various components of plaques. A, Type I-II lesion in common carotid artery (layered foam cells were detected by histology). On multicontrast-weighted MR images, the carotid wall appears normal (arrow). The asterisk indicates the lumen; H&E, hematoxylin and eosin. B, Type III lesion in common carotid artery (kipid-rich necrotic core was detected by histology). On MR images, carotid wall shows slight eccentric thickening (arrow). C, Type IV-V lesion in internal carotid artery (lipid-rich necrotic core was detected by histology). On MR images, lipid-rich necrotic core (arrow) had iso-signal intensity on both T1WI and TOF but iso-signal intensity to slightly high signal intensity on PDWI and T2WI. Lumen is moderately stenosed. D, Type VI lesion just distal to carotid bifurcation (acute to subacute mixed hemorrhages were detected by histology). On MR images, acute and subacute mixed hemorrhage had high signal intensity on both TOF and T1WI, iso-signal intensity to slightly high signal intensity on PDWI and T2WI (arrow). E, Type VII lesion in common carotid artery. Extensive calcification was present in plaque by histology (boundary of calcified region is outlined for clarification). Calcified region (arrow) had low signal intensity on all images. F, Type VIII lesion in internal carotid artery. Connective tissue was characterized by immunocytochemistry as proteoglycan-rich early matrix. MR images show varied signal intensity. PDWI, proton density-weighted imaging; T1WI, TI-weighted imaging; T2WI, T2-weighted imaging; T0F, time of flight. (Modified from Cai JM, Hatsukami T, Ferguson MS, et al: Classification of human atherosclerotic lesions with in vivo multicontrast magnetic resonance imaging. Circulation 106:1368, 2002.)

Vulnerable plaque refers to thrombosis-prone plaques and plaques with high probability of undergoing rapid progression and thus becoming culprit plaques. 11 The salient characteristics of vulnerable and stable plaques are listed in Table 28-4. 12

Large Lipid-Rich Necrotic Core. The large lipid-rich necrotic core is composed of extracellular lipids (cholesteryl

Vulnerable Plaque



TABLE 28-3	MRI Characteristics of Various Tissue Components in Different MRI Sequences					
	T1	T2	Proton Density	Enhancement		
Lipid	Hyper	Нуро	Hyper	None		
Fibrocellular	Hyper	Hyper	Hyper	None		
Calcium	Нуро	Нуро	Нуро	None		
Necrotic core	Low	High	High	None		
Fibrous cap	Low	Low	Low	Enhancement		
Hemorrhage	High	Low	Intermediate	None		

TABLE 28-4

Characteristic Features of Vulnerable and Stable **Plaques**

Large lipid-rich core (> 40% plaque volume) Thin fibrous head (< 65-100 gm) Active inflammation Endothelial denudation with superficial platelet aggregation

Fissured plaque Stenosis > 90%

Superficial calcified nodule Glistening yellow (large lipid-rich core, thin cap) Intraplaque hemorrhage Endothelial dysfunction Outward remodeling

Stable Plaque

Small lipid core Thick fibrous cap Abundant smooth muscle cells No inflammation

monohydrate or unesterified cholesterol) and cellular debris. Necrotic core has low intensity on T2-weighted images because of short T2 and no contrast enhancement.

Thin Fibrous Cap. The fibrous cap is made up of intimal smooth muscle cells and connective tissue. Fibrous tissue has short T1, which results in bright signal on T1-weighted images and contrast enhancement. Rupture-prone plagues are very thin (65 to 150 g m), which is below the spatial resolution of MRI. In three-dimensional time of flight, the fibrous cap is seen at the interface of the bright lumen and dark wall, 13 making it possible to assess its integrity (Fig. 28-2). ¹⁴ Higher strength magnets and multiple-channel coils may be necessary for further characterization. Easy visualization of a head on MRI generally indicates that it is relatively thick. Nonvisualization (implying a thin cap) or disruption of the fibrous cap indicates potentially high risk plaques (Fig. 28-3).

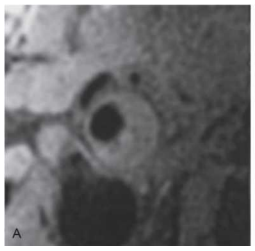
Active Inflammation. Inflamed plaques are characterized by rich and active macrophages, mast cells, T cells, proinflammatory cytokines, procoagulant mediators, increased matrix metalloproteinase expression, and reduced smooth muscle cells. Inflammation can be detected by contrast enhancement because of increased vascular permeability and edema 9 or by molecular MRI techniques (see later).

Erosion or Fissure of Plaque Surface. Ulcerations are more common in symptomatic patients, regardless of the intensity of symptoms. In addition to the multicontrast black blood MRI, longitudinal black blood magnetic resonance angiography (MRA) increases the ability to identify ulcerations.

Superimposed Thrombus. Thrombus is more prevalent in symptomatic patients, especially with ipsilateral symptoms and plaque ulceration. Thrombotically active plaque associated with high inflammatory infiltrate is seen in 74% of patients presenting with major stroke. Thrombus is the plaque component that is more heterogeneous and difficult to detect.

Luminal Stenosis > 90%. Shear stress imposes a significant risk of thrombosis and sudden occlusion. It also indicates the presence of many nonstenotic or less stenotic plaques that are vulnerable to rupture and thrombosis.

Superficial Calcified Nodules. Calcified nodules within or close to the head can protrude through and rupture the head. Surface calcified nodules can have an exposed thrombogenic surface or can be encapsulated. Calcification (mainly calcium hydroxyapatite) has a low signal because of low proton density and diffusion-mediated susceptibility effects, making



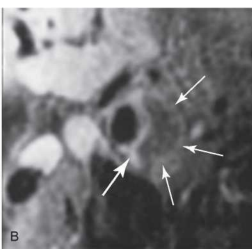


FIGURE 28-2 Axial precontrast (A) and postcontrast (B) double inversion recovery fast spin-echo black blood images of the carotid artery with fat saturation show heterogeneous enhancement along the outer wall of the atheroma (arrows) and along the margin of the lumen, indicating a fibrous cap. (Modified from Wasserman BA, Smith WI, Trout HH, et al: Carotid atherosclerosis: in vivo morphologic characterization with gadolinium enhanced double oblique MR imaging. Initial results. Radiology 223:566, 2002.)

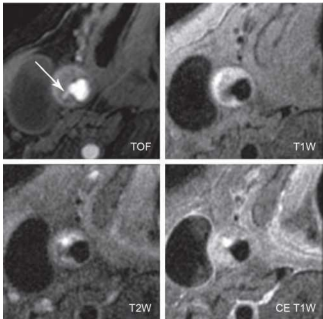


FIGURE 28-3 Multicontrast MR images demonstrating rupture of fibrous cap in a carotid artery plaque (arrow). High signal intensity in T2-weighted and postcontrast TI-weighted images indicates plaque inflammation and neovasculature. (Modified from Chu B, Ferguson MS, Chen H, et al: Cardiac magnetic resonance features of the disruption-prone and the disrupted carotid plaque. JACC Cardiovasc Imaging 2:883, 2009.)

it is difficult to be detected in black blood imaging, but it can be seen clearly in time-of-flight images.

Intraplaque Hemorrhage. Intraplaque hemorrhage is due to rupture of fragile microvessels. It has a variable signal, depending on the blood breakdown products. Methemoglobin , which is seen in the subacute stage, shows T1 shortening and high signal on T1 imaging. The window for observing these changes is small, and it is essential to localize the hemorrhage within the plaque. Hemorrhages close to the lumen and intraluminal thrombosis are more often associated with embolism and symptoms than deep plaque hemorrhages are.

Outward Positive Remodeling. Vulnerable plaques are often associated with positive remodeling, which prevents luminal narrowing; as a result, the patient may be asymptomatic, and stress perfusion imaging is often normal. ¹⁵ This is a surrogate marker of plaque vulnerability.

Adventitial Inflammation and Neovascularity. Adventitial inflammation and neovascularization are associated with high risk of plaque rupture. Gadolinium enhancement in T1-weighted MRI has high sensitivity for neovascularization (80%). Dynamic enhancement indicates vascular density, which is a marker of neovascularity, and delayed enhancement reflects increased permeability of inflamed vessels. ¹⁶

Post—Gadolinium Contrast Enhancement

Intense contrast enhancement indicates neovascularity and increased endothelial permeability, which results in entry of the contrast agent from blood plasma into the plaque. Strong enhancement is an indicator of plaque inflammation, which is a sign of vulnerability. The amount of enhancement is a marker of the extent of acute inflammation. Late gadolinium enhancement is used to identify fibrous plaque. ^{13,17}

Plaque Quantification

Plaque seen on black blood MRI has been validated with transesophageal echocardiography ¹⁸ and with histopathology in animal models in multiple studies. Plaque can be measured manually or by automated techniques, such as MEPPS

(morphology-enhanced probabilistic plaque segmentation algorithm) and CASCADE (computer-aided system for cardiovascular disease evaluation), which are based on probabilistic assumptions. The system determines the probability that each MRI pixel belongs to each of the four tissue types, namely, lipid, calcification, fibrous tissue, and loose matrix, and uses computing active contours to identify the boundary lines of high probability regions of each tissue, which are then segmented and displayed in three dimensions. ¹³ This reduces analysis time and reader bias and improves reproducibility. Quantitative measurements have been correlated with histology and are highly reproducible. ¹³

MRI of Individual Vascular Territories

The majority of the work on morphological MRI of the atherosclerotic plaque has been performed in the carotid arteries and aorta because of their superficial location and large size, respectively. New studies are reporting assessment of plaques in coronary and other arterial territories, which are limited by small size and deep location. In all these territories, MRI acquisition takes a long time because of the need for high-resolution images.

Carotid Plaques. Black blood multicontrast MRI with surface coils has been well established and validated with histology in the demonstration of carotid plaque morphology (volume, area, and thickness), structure, and composition. ¹⁹ There is a correlation between thinning or rupture of the fibrous cap and recent history of transient ischemic attack or stroke. ²⁰ Ipsilateral arteries in symptomatic patients have a higher prevalence of intraplaque hemorrhage and fibrous cap rupture. Intraplaque hemorrhage indicates an increase in plaque burden (increased necrotic core size), decreased lumen size, repeated intraplaque hemorrhage, and subsequent plaque rupture, making it a superior prognostic indicator (Fig. 28-4).

A greater plaque burden and plaque eccentricity is prevalent among patients with prior major cardiovascular or -cerebrovascular events. ²¹ A prospective study of asymptomatic patients with a 50% to 79% stenosis showed that intraplaque hemorrhage, larger necrotic core area, and thinned or ruptured fibrous cap are associated with a high hazard ratio for subsequent clinical events (Fig. 28-5). Increases in thickness of arterial wall, average area of intraplaque hemorrhage, and necrotic core area were associated with increased hazard ratio for subsequent clinical areas. ²² MRI was also useful for evaluation of plaque regression in carotids (eg, ORION [Outcome of Rosuvastatin treatment on carotid artery atheroma: a magnetic resonance Imaging Observation] trial).

Aortic Plaque. Multiple studies have established a correlation between aortic plaques and coronary artery lesions. Thoracic aortic plaques are associated with hypercholesterolemia , and a correlation with low-density lipoprotein cholesterol (LDL-C) level has been shown. ⁵ Abdominal aortic plaques are not associated with LDL-C levels, but they are associated with smoking. Although the exact reason for this variable susceptibility to different risk factors is not known, there are slight differences in the vascular beds. The abdominal aorta has less prominent vasa vasorum, resulting in nutrition only by diffusion, and it has higher blood pressures and higher stiffness with more collagen and less elastin and geometric tapering. Plaques in both the thoracic and abdominal aorta are associated with age and high blood pressure. ¹⁸

There is a high association between thoracic aortic plaques and coronary artery disease (CAD), more than with carotid or femoral artery plaques. Complex plaques are associated with cardiovascular events. Autopsy studies have shown a relationship between severe abdominal aortic plaques and cardiac events, but no association has been shown between CAD and abdominal plaques. An MRI study ¹⁸ showed a higher

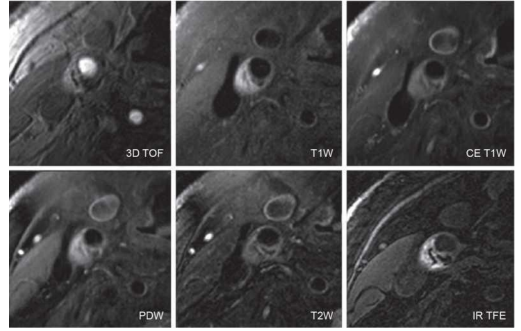


FIGURE 28-4 Hyperintense signal in the plaque is due to intraplaque hemorrhage on the three-dimensional time-of-flight (3D TOF), T1-weighted (T1W), and inversion recovery turbo field-echo (IR TFE) sequences. Hypointense signals in both TOF and IR TFE techniques are the densely calcified areas. (Modified from Chu B, Ferguson MS, Chen H, et al: Cardiac magnetic resonance features of the disruption-prone and the disrupted carotid plaque. JACC Cardiovasc Imaging 2:883, 2009.)

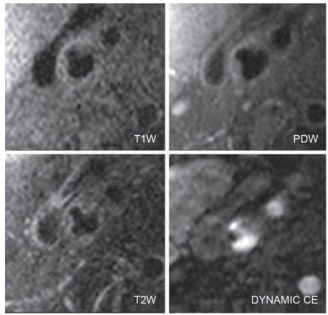


FIGURE 28-5 Multicontrast MR images show a large surface ulceration in a carotid artery plaque. (Modified from Chu B, Ferguson MS, Chen H, et al: Cardiac magnetic resonance features of the disruption-prone and the disrupted carotid plaque. JACC Cardiovasc Imaging 2:883, 2009.)

prevalence of plaques in the thoracic (73%) and abdominal aorta (94%) in patients with CAD than in those without it. The extent of plaques correlated with the extent of coronary artery stenosis. Whereas the thoracic plaques were independently associated with CAD, the abdominal aortic plaques were not independently associated with CAD. There was no difference in prevalence between CAD with and without MI.

Complex plaques in the abdominal aorta were more slow in CAD patients with MI than without MI. Among patients without

MI, complex aortic plaques were more prevalent in patients with complex coronary lesions than without complex coronary lesions. All these suggest that complex aortic plaques are linked to coronary plaque instability, which leads to the development of MI and complex coronary lesions. Complex aortic plaque, especially in the abdominal aorta, is a good marker of coronary plaque instability. CT studies have also shown a better association between thoracic aortic plaques and CAD. Hence, thoracic aortic plaque is a better marker of coexisting CAD. ⁵

Coronary Artery Plaque. Plaque imaging in coronary arteries is hampered by the small diameter, tortuosity, deep location, and high mobility of the coronary arteries and the small volume of plaque. In addition, not all ruptured plaques cause acute coronary syndromes, and vulnerability is not static. A vulnerable plaque one day may look stable another day. ²³ There has been a good correlation between black blood MRI-measured coronary wall thickness and matched pathology sections, with good reproducibility. Increased wall thickness is seen in patients with more than 40% stenosis on angiography. ²⁴ Because of low resolution, MRI may overesti mate coronary wall thickness in normal patients.

In asymptomatic type 1 diabetes mellitus, coronary wall MRI reveals higher plaque burden in subjects with nephropa thy than with normoalbuminuria. ²⁵ Hyperintense plaque in T1-weighted sequences indicates an unstable plaque due to intraplaque hemorrhage, and it correlates with positive remodeling, ultrasound attenuation, lower Hounsfield units in CT, and transient flow after percutaneous coronary intervention. These findings are similar to those in carotid plaque characterization, which has been validated with histology. ²⁶ T1-weighted coronary plaque imaging has the potential for identification of complex coronary lesions in patients with unstable CAD.

High-resolution coronary plaque imaging has been performed in humans by 3T scanners with a 32-channel cardiac coil (0.5 x 0.5 x 3-mm resolution) and in animals by a 9T scanner (97- q m resolution) and intravascular MRI (117 x

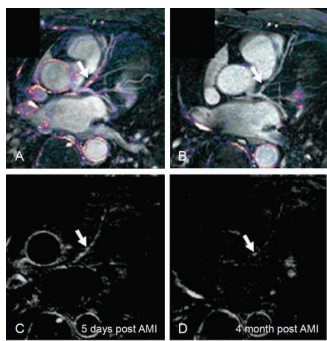


FIGURE 28-6 Coronary MRA of a 62-year-old patient with inferolateral infarction shows a luminal stenosis in the proximal left anterior descending (LAD) coronary artery (arrow) that corresponds to 50% stenosis in x-ray angiography (A) . MRA 5 days after infarction (C) shows contrast uptake within proximal LAD, MRA 4 months after MI (B. D) shows marked reduction of coronary enhancement in proximal LAD; the stenotic region shows mild uptake (arrow). A and B have fused images, which fused the enhancement in C and D with MRA. (Modified from Ibrahim T, Makowski M, Jankauskas A, et al: Serial contrast-enhanced cardiac MRI demonstrates regression of hyperenhancement within the coronary artery wall in patients after acute myocardial infarction. JACC Cardiovasc Imaging 2:580, 2009.)

156- im resolution). 27 A study demonstrated coronary arterial wall enhancement during acute MI due to edema and inflammation and resolution during delayed phase (6 months later), which offers the potential for visualization of inflammation in coronary atherosclerosis (Fig. 28 - 6). 28,29 However, enhancement was diffuse and not localized to the occlusive site.

MRI in Plaque Progression and Regression

Because of its high reproducibility, MRI is useful in the evaluation of plaque progression and regression and has been well established in numerous animal and human studies. The use of MRI has significantly reduced the sample size required for evaluation of pharmacological regression of atherosclerosis in clinical trials. In an early study, 30 balloon-injured rabbits fed a high- versus low-cholesterol diet, MRI at 4 and 20 months showed higher wall thickness and stenosis in the highcholesterol similar MRI group. Α study with hypercholesterolemic rabbits fed a high-cholesterol diet for 12 weeks followed by a normal diet for 12 weeks showed increased wall volume during a high-cholesterol diet phase and decreased wall volume during the normal diet phase. 31

Numerous MRI studies have evaluated the response of carotid and aortic atherosclerotic lesions to various doses and regimens of medical therapy. ⁵ MRI can measure plaque reduction and map changes in plaque composition in response to intensive lipid-lowering treatment and has been validated with histology. MRI has good reproducibility and low inter scan variability and is able to detect a 10% change in wall volume and a 20% change in percentage of lipid-rich or necrotic core components. 32

In a study of asymptomatic subjects with 50% carotid stenosis, MRI showed that the rate of increase in carotid wall areas at 18 months was lower in those who took statins compared

with the non-statin group. 33 In a similar study of CAD patients, MRI showed significantly less lipid core area and lipid composition in patients who were receiving intensive lipidlowering therapy compared with a nontreated group. 34 MRI of the thoracic aorta demonstrated changes in plaque volume and lumen after simvastatin (20 to 80 mg) therapy for 6 months that correlated with LDL reduction. 35 Other studies have shown a continued decrease in plaques in the aorta and carotid arteries at 12 months and after 18 to 24 months of statin treatment, ³⁶ with minimal changes in luminal dimension.

A study comparing 5 mg versus 20 mg of atorvastatin for 12 months in hypercholesterolemic patients showed a greater reduction of thoracic aortic plaque in the 20-mg than in the 5-mg group that correlated with reduced LDL-C levels. Abdominal aortic plaque did not change with the 20-mg dose but became 28 worse with just 5 mg ³⁷ (Fig. 28-7). Similar results were shown in a Japanese study that used 20 mg of atorvas tatin. Another study that compared 20 mg versus 80 mg of simvastatin showed greater reduction of carotid and aortic plaque at 12 months in the higher dose group. 38 All these studies prove that intensive atorvastatin therapy reduces LDL-C and causes plaque regression in the thoracic aorta and carotid and coronary arteries.

Fibrates reduce triglycerides and increase high-density lipoprotein cholesterol (HDL-C) levels. An MRI study of hypertriglyceridemic patients after 400 mg of bezafibrate showed regression of thoracic aortic wall plaques without change in cross-sectional area, regression of abdominal aortic plaques with increase in cross-sectional area, and correlation between wall area change and triglyceride reduction and HDL-C increase. 39 The change in abdominal plaques alone correlated with HDL particle size reduction and LDL size increase by nuclear magnetic resonance, implying that there might be different mechanisms in the thoracic and abdominal aorta for plaque regression and that triglyceride plays an important role in atherosclerosis in the abdominal aorta.

Because the abdominal aorta has no vasa vasorum and reverse lipid transport happens through diffusion into the lumen, there might be an increase in area after therapy, compared with the thoracic aorta, in which reverse transport occurs because of the presence of vasa vasorum and there is no significant reduction. ¹⁸ MRI has also been used to compare the effects of statins versus statins plus the peroxisome proliferator activated receptor y agonist on plaque. 40

Advances in Morphological MRI

Higher field strength of magnets, dedicated multichannel phase array coils to increase signal-to-noise ratio, multislice motionsensitized driven-equilibrium turbo spin-echo sequences to suppress plaque-mimicking artifacts, and three dimensional isotropic sequences for evaluation of luminal surface and plaque definition are some of the latest advances in morphological MRI. T2* imaging of iron forms within the plaque has demonstrated lower T2* values in symptomatic plaques. 41

3T MRI has the advantages of having twice the signal of 1.5T MRI and higher contrast-to-noise ratio, although it has higher chemical shift and susceptibility artifacts. Double inversion delay time is increased to accommodate increased blood T1. High spatial resolution images are acquired by parallel imaging and multiple averages, and these are useful for detection characterization, and quantification of early and moderate plaques, with a high correlation between plaque quantity and components and histopathology. An 18-month follow-up study using 3T MRI showed 50% reduction in lipid content with minimal change in plaque size in a subject receiving

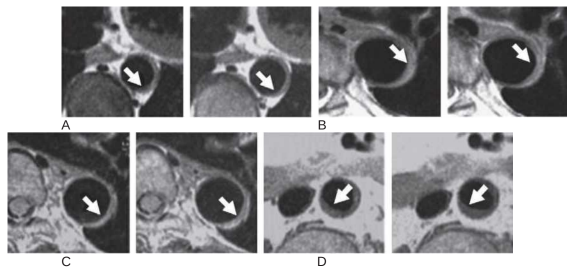


FIGURE 28-7 MRI at baseline and after 1 year of atorvastatin treatment. In **A**, a thoracic aortic plaque showed 28% vessel wall area reduction with a 20-mg dose of atorvastatin. In **B**, a thoracic aortic plaque showed 15% progression with a 5-mg dose. In **C**, a thoracic aortic plaque showed 15% progression with a 5-mg dose. In **D**, an abdominal plaque showed 9% progression despite a 20-mg dose. (Modified from Yonemura A, Momiyama Y, Fayad ZA, et al: Effect of lipid lowering therapy with atorvastatin on atherosclerotic aortic plaques detected by noninvasive MRI. J Am Coll Cardiol 45:733, 2005.)

aggressive statin therapy. Similar studies have been performed in coronary arteries, thoracic aorta, and femoral artery ies. Higher field strengths, such as 7T and 9.4T, increase spatial resolution further and suggest potential for spectroscopic methods. 42

Coronary Magnetic Resonance Angiography

Coronary MRA provides information on the coronary artery lumen and vessel wall including plaque, which can be threedimensionally reconstructed in multiple planes, without radiation or intravenous administration of contrast material. However, valuation of coronary arteries requires high spatial resolution (3 to 4 mm) and temporal resolution (< 75 msec). A three-dimensional steady-state free precession whole-heart sequence with T2 preparation to decrease myocardial signal and fat saturation to decrease epicardial fat signal is used to image coronary arteries. Blood vessels appear bright because of intrinsic contrast. Images are acquired in diastole and expiration with free breathing technique using navigator gating of the diaphragm (Fig. 28-8). To obtain high-resolution images (0.7 to 0.8 mm), acquisition time can be very long, up to 12 to 15 minutes. MRI has a sensitivity of 80% to 90%, specificity of > 90%, and negative predictive value of 81% for identification of coronary artery stenosis. 43 However, coronary MRA is not as widely used as CT angiography because of low sensitivity, long acquisition time, and lack of expertise in many hospitals.

Magnetic Resonance Angiography

Whole-body MRA can examine the entire arterial tree excluding the intracranial and coronary vessels in one sitting by use of bolus chase technique, multichannel receiver surface coils, and parallel imaging techniques, without radiation or arterial cannulation, and it is less nephrotoxic. Multistation techniques are used, typically a four-station technique; the first station covers supra-aortic arteries and the thoracic aorta, the second station covers the abdominal aorta, the third station



FIGURE 28-8 Three-dimensional whole-heart steady-state free precession MRI sequence demonstrates the origins and proximal segments of all the left and right coronary arteries.

covers the external iliac to popliteal areas, and the fourth station covers up to the ankle. Subsystolic venous compression in the calf can reduce venous contamination and increase signal-to-noise and contrast-to-noise ratios. Higher resolution scans can be performed at sites of plaques to characterize them and to quantify luminal stenosis. 3T magnet (higher signal), surface multichannel coils, parallel imaging, and blood pool agent (MS-325) will potentially improve the quality of MRA. MRA has been validated for pelvic and lower limb arteries. However, reproducibility has not been completely assessed yet, and it can overestimate stenosis.

The ability to image multiple vascular beds makes it useful in the evaluation of atherosclerosis. MRA can be combined with

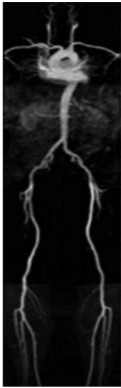


FIGURE 28-9 Maximum intensity projection of whole-body MRA shows all the arteries from the head to the leg, which are used to calculate the atherosclerotic score index. (Modified from Lehrke S. Egenlauf B. Steen H. et al: Prediction of coronary artery disease by a systemic atherosclerosis score index derived from whole-body MRI angiography. J Cardiovasc Magn Reson 11:36, 2009.)

whole-body MR screening. 44 MRA has detected unknown arterial disease in different populations, such as those characterized by advanced age, diabetes, and post-MI. 45 MRA is useful for repeated clinical examinations but has not yet been studied or validated to predict future cardiovascular events. An atherosclerotic score index (ASI) has been measured by scoring luminal narrowing (1, normal; 2, < 25%; 3, 26% to 50%; 4, 51% to 75%; 5, 76% to 99%; 6, occlusion) in 40 extracardiac segments and dividing the sum by the number of analyzable segments (Fig. 28-9). ASI was higher in patients with CAD than in those without and correlated with Framingham and PROCAM risk scores. An ASI > 1.54 has 59% sensitivity, 86% specificity, and 84% positive predictive value for predicting CAD and confirming increased risk of CAD in those with extracardiac atherosclerosis, in addition to giving an estimate of total atherosclerotic burden in the body. 45

Peripheral MRI

Peripheral arterial atherosclerosis is the most common cause of peripheral arterial disease, often leading to vascular obstruction. MRA, high-resolution black blood imaging, and first-pass contrast-enhanced dual contrast perfusion imaging of the calf are the MRI techniques used in the assessment of peripheral arterial disease. MRA can detect and quantify luminal stenosis, although it is technically challenging because of artifacts such as signal loss from in-plane saturation, turbulent flow, metallic clips and stents, and end-organ effects of peripheral arterial disease. Highresolution black blood imaging using surface coil and flow saturation can diagnose preclinical vascular disease, measure plaque volume, assess disease severity, and monitor plaque progression or regression in response to therapy.

The superficial femoral artery is the ideal vessel because it is

Myocardial Function

particularly at maximal workload. 47

Global Myocardial Function

Left ventricular volumes, mass, and ejection fraction are important markers of cardiovascular disease and independent - 28 predictors of cardiovascular events, with changes seen before the onset of symptoms. Left ventricular hypertrophy is considered an abnormal response to conditions such as hypertension, not physiological or compensatory, and it is an independent predictor of cardiovascular events. 23 Left ventricular mass is an important marker of subclinical disease and decreases after therapy reduces adverse events. 23 The MESA study has demonstrated an 18-fold higher rate of congestive heart failure during a short period of observation in those with left ventricular hypertrophy, after accounting for traditional risk factors and coronary calcium score. 43

superficial and nonmobile. Popliteal arteries have also been evaluated to demonstrate the changes that occur with remodeling and restenosis after angioplasty. 46 First-pass contrast-enhanced dual contrast perfusion imaging of calf muscle at peak exercise with an MR-compatible pedal ergometer and ³¹ P MR spectroscopy at peak exercise measure phosphocreatine recovery kinetics, which varies greatly between normals and patients with peripheral arterial disease,

MRI is the most accurate technique in the evaluation of cardiac volumes and mass, with only 5% standard errors compared with 20% for echocardiography. 48 Volumes are measured manually or semiautomatically by steady-state free precession sequences obtained in the short-axis plane, by drawing manual or automatic endocardial and epicardial contours. Mass is obtained by multiplying the wall volume with myocardial density (1.05 g/cm³) (Fig. 28-10). With increasing age, particularly in men, the mass-to-volume ratio increases, although left ventricular mass is maintained because of a decrease in end-diastolic volume due to concentric remodeling.

Regional Myocardial Function

Regional changes often precede global abnormalities in cardiac function. Strain is the fractional change in length from the resting state in diastole to contraction in systole. Radial, circumferential, and longitudinal strains can be measured on MRI by myocardial tagging techniques, such as SPAMM (spatial modulation of CSPAMM magnetization) or (complementary modulation of magnetization), which produces dark saturation bands perpendicular to the scanning plane ⁴³ (Fig. 28-11). Strain is quantified by harmonic phase analysis, which is more accurate than visual estimation of regional function. Strains can be measured separately in the subendocardial, mid-wall, and subepicardial layers. Other techniques for tagging are strainencoded imaging (tags are parallel to imaging plane), displacement encoding with stimulated echoes (DENSE), and velocity encoding phase shifts with phase contrast imaging.

With age, although the ejection fraction is maintained in the normal range, peak strain decreases, particularly in the left anterior descending territory of men. In women, strain increases with concentric remodeling in all but the highest quartile of mass-to-volume ratio. A direct relationship between regional diastolic dysfunction (decreased diastolic strain) and increasing left ventricular mass is demonstrated in asymptomatic individuals. After controlling for established risk factors, increased diastolic blood pressure was associated

RESULTS SUMMARY:

Ejection fraction 37.7% Stroke volume 56.4 mL Stroke index n/a Cardiac output: 3.4 1/min Cardiac index: n/a

ED time ED volumes ES time ES volumes ED wall mass ED wall + papillary mass ED wall + papillary	0.00 ms (phase 1) 149.39 mL 376.00 ms (phase 12) 93.00 mL 81.47 9 N/A N/A
ED wall + papillary - correct. mass Heart rate Patient height Patient weight	60.00 bpm n/a 80.00 kg

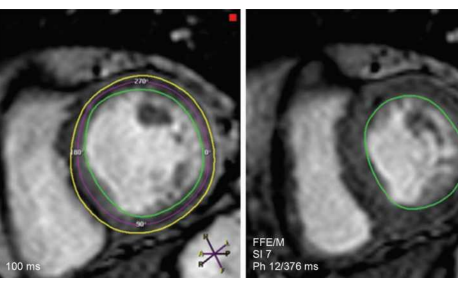


FIGURE 28-10 MRI measurement of left ventricular volumes, function, and mass. Endocardial (green) and epicardial (yellow) contours are shown.

with decreased circumferential strain in asymptomatic individuals, 43 particularly in smokers.

Stress Perfusion MRI

First-pass myocardial perfusion imaging with MRI is a sensitive technique to detect myocardial ischemia, with high detection of CAD compared with coronary angiography, 49 and it has a 100% negative predictive value for subsequent CAD. 50 Perfusion MRI is performed both at rest and after pharmacological stress (adenosine or dipyridamole). Intravenous contrast produces T1 shortening of normal myocardium, resulting in bright signal on T1-weighted sequences. Hypoperfused areas (ischemic or revascularized infarction) appear dark (Fig. 28-12). Microvascular dysfunction is seen as a subendocardial perfusion defect in a nonvascular distribution. Sequences used for perfusion are typically multislice T1-weighted two-dimensional sequences, which could be steady-state free precession (SSFP), fast low-angle shot (FLASH), or gradient-recalled echo planar imaging (GRE EPI), either inversion or saturation recovery based or hybrid, each with its advantages and disadvantages. 51 Typically, perfusion studies are followed by delayed enhancement images to evaluate myocardial viability, 52 and this enhances the performance of adenosine stress cardiovascular magnetic resonance. Detection of even small amounts of myocardial delayed enhancement in patients without known MI is an adverse prognostic indicator. Direct assessment of myocardial perfusion after adenosine stress is sensitive, whereas adenosine-induced wall motion abnormalities are highly specific. ⁵³

Semiquantitative analysis of the blood flow could be obtained by analysis of the postcontrast myocardial signal intensity as a function of time. Myocardial perfusion reserve index is the ratio of the maximal upslope of the time-intensity profiles at stress and at rest, which is a good indicator of perfusion. After vasodilation in a normal vessel, there is a peak followed by washout before plateau. If there is no over shoot, it indicates no vasodilator response. Absolute quantification is possible if there is an accurate estimate of arterial input function, which is measured from the left ventricular



250

200

150 100 50 ED

ES

FIGURE 28-11 Regional myocardial function evaluated by grid tags with SPAMM technique. Endocardial and epicardial contours are drawn to estimate strain at multiple levels (bottom left). Strain map is illustrated in the bottom right image.

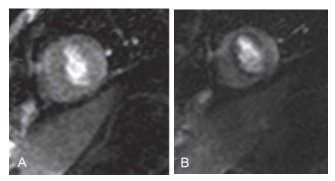


FIGURE 28-12 MRI in a 46-year-old woman presenting with exertional chest pain shows normal rest perfusion scan (A), but there is a dark band of hypoperfusion in the apical septal, anterior, and inferior walls in the stress image (B) consistent with ischemia in the distal left anterior descending territory.

blood pool signal. Dual bolus or dual sequence methods improve accuracy. Quantification of myocardial blood flow and myocardial perfusion reserve can be performed through deconvolution models or model-independent analysis. MRI has a higher accuracy for subendocardial than for transmural perfusion analysis. ⁵⁴

Perfusion imaging can also be performed without intrave nous contrast material by endogenous contrast mechanisms. In the BOLD (blood oxygen level dependent) technique, nor mally perfused areas have oxyhemoglobin, which is slightly diamagnetic, producing normal MRI signal, but hypoperfused areas have deoxyhemoglobin, which is paramagnetic and causes signal loss in T2*/T2-weighted images. ⁵⁵ ASL (arterial spin labeling) works by magnetically labeling blood flowing into slices of interest, which then exchanges with tissue water, altering the tissue magnetization. Subtraction of the image with labeled inflowing spins from an image without spin labeling gives a perfusion-weighted image.

Dobutamine MRI

Dobutamine infusion makes ischemic segments dysfunctional, which could be assessed by cardiac MRI as an alternative test for the diagnosis of CAD and quantifying myocardium at risk, particularly useful in patients in whom echocardiography cannot

be performed. The sensitivity of dobutamine MRI is higher than that of echocardiography, ⁵⁶ and it can be further enhanced by myocardial tagging. ⁵⁷ It is also useful in predicting left ventricular functional recovery after coronary revascularization. ⁵⁸ Combined adenosine and dobutamine stress imaging showed 99% survival in patients with normal images and 84% with abnormal images in a series of 513 patients with known or suspected coronary disease during a mean follow-up of 2.3 years. ⁵⁹ In a multi center, multivendor trial, perfusion cardiovascular magnetic resonance was proven to be a valuable alternative to SPECT for CAD detection, showing equal performance in the head-to-head comparison. ⁴⁹ It is also useful in assessing the results of a percutaneous coronary intervention.

Intravascular MRI

Intravascular MRI (IVMR) can be used to image small vessels, such as coronary arteries and iliac or renal arteries, with high signal-to-noise-ratio, spatial resolution, and temporal resolution. ⁶⁰ Intravascular coils have been used for lesion assessment and characterization in carotid and iliac arteries, although there was not enough spatial resolution to visualize thin fibrous cap or atheroma thickness. IVMR can also evaluate increased lipid levels associated with potentially vulnerable plaques, with the help of an integrated, self-contained MRI probe, with magnets and transmit-receive radio frequency coils on the tip of the catheter that measure lipid concentration in the arterial wall based on apparent diffusion coefficient (ADC). Fibrous tissue has high ADC (unrestricted diffusion), but lipid-rich tissue has low ADC (restricted because of large cholesterol esters).

Intravascular coils within the venous system can evaluate the adjacent arterial system. IVMR is potentially useful for guiding coronary interventions such as angioplasty and stent placement, with good real-time visualization of the catheter and three-dimensional multiplanar reconstruction capabilities and without need for fluoroscopy, radiation, or nephro toxic contrast agents. IVMR is also used to confirm delivery of local gene therapy or nanoparticles to plaques or vascular wall by a special balloon catheter system. Limitations of intravascular systems are their invasiveness, catheter size, length of imaging protocols, and local heating.

Shear Stress

Plaque rupture is frequently seen in areas exposed to high wall shear stress, typically upstream of maximal stenosis. High wall shear stress also induces antiproliferative activity by endothelial cells on the upstream side, which enhances plaque vulnerability. Information on vascular geometry, inflow conditions, and plaque is acquired from MRI including angiography, and a computational fluid dynamic model is created from which shear stress acting on the luminal surface can be computed from the velocity field. It is unclear how variable geometric reconstruction and restricting assumptions on blood rheology or vessel wall compliance affect the accuracy of results. ¹³ Wall shear stress can also be calculated semiautomatically and with good reproducibility with model-based segmentation of phase contrast MRI by determination of flow volume and maximal flow velocity in cross sections of these vessels (Fig. 28-13). There is site specific variation of wall shear stress based on this method.

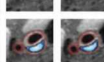
Vascular Function

Vascular function is an early marker of cardiovascular disease. Arterial compliance is both a cause and a consequence of











■ Lipid/necrotic core O Intraplaque hemorrhage ■ Ulcer

Follow-up

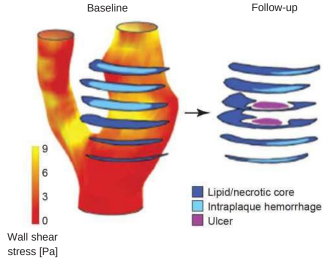


FIGURE 28-13 Morphologic MR images and vessel wall segmentation at baseline and 10month follow-up. Initial scan shows lipid-rich necrotic core (blue) and intraplaque hemorrhage (teal) in internal carotid artery. Follow-up MRI shows large ulcer in the proximal internal carotid artery. Ulceration developed in the region that had the highest wall shear stress at baseline. (Modified from Chu B, Ferguson MS, Chen H, et al: Cardiac magnetic resonance features of the disruption-prone and the disrupted carotid plaque. JACC Cardiovasc Imaging 2:883, 2009.)

vascular disease. Aortic stiffness is associated with cardiovascular risk factors, morbidity, and mortality. It has been shown to be independently associated with CAD. In addition, decreased central aortic compliance causes systolic hypertension left ventricular hypertrophy, and diastolic dysfunction. Velocity-encoded phase contrast MRI is a useful noninvasive technique in the assessment of various vascular parameters compared with invasive techniques, which often make assumptions on central arterial pressure and the path length. 61 Various vascular parameters could be assessed by MRI (Table 28-

Molecular Imaging

Molecular imaging elucidates the biology of physiologically relevant cellular and molecular targets in atherosclerosis by use of specific contrast agents, which enables detection before

the development of gross phenotypic changes. Molecular imaging complements anatomical and physiological imaging and can be performed with nuclear medical techniques such as SPECT or PET, MRI, CT, ultrasound, and optical imaging, either as stand-alone or hybrid modalities. 62 Although many of these techniques are still in basic and translational research stages, recent advances in understanding of pathophysiology, imaging agent chemistry, and imaging platforms have led to the progression of many agents to clinical evaluation and application , which includes therapy. 63

TABLE 28-5	Vascular F MRI	unctional Parameters That Could Be Assessed by
Parameter		Measurement
Aortic strain		(systolic diameter — diastolic diameter)/ diastolic diameter
Aortic stiffness in	dex (p)	In(systolic blood pressure/diastolic blood pressure)/[(systolic diameter — diastolic diameter)/diastolic diameter]
Aortic distensibility		
		2 X (systolic diameter — diastolic diameter)/[systolic blood pressure — diastolic blood pressure) X diastolic diameter]
Aortic elastic modul	us E _P	(systolic blood pressure — diastolic blood pressure)/[(systolic diameter — diastolic diameter)/diastolic diameter]
Young's circumferer static elastic mo		E _P X diastolic diameter/2h, where h is the diastolic intima-media thickness
Pulse wave velocity	(m/sec)	Distance between two points/transit time of pulse wave between points

Molecular imaging provides insights into pathophysiology that reveal disease diversity. Its advantages include early and refined diagnosis, identification of vulnerable plaque and highrisk patients, and multimodality imaging options. Initiation and titration of therapy targets the biology and molecular profile of vulnerable plaque rather than the lipid profile. It provides opportunities for image-guided therapy and local drug release, and systemic toxicity is decreased. Immediate and accurate follow-up is possible because the molecular response to therapy is quantified. Surrogate imaging endpoints for clinical trials enable assessment of the efficacy of novel drugs that are faster and cheaper.

Components of Molecular Imaging

Molecular imaging involves target molecules, ligands, carrier vehicles, and signal elements (Fig. 28-14). Although there are various possible target molecules for atherosclerosis, imaging of them is challenging because of their location in vessels deep inside the body, small quantities, motion, and high shear stress in large arteries. 64 Ligands are molecules such as monoclonal antibodies, fragments, small molecules, small peptides, or carbohydrates that have high affinity and specificity for the target molecule. Probes can also be modified to be taken by specific cells. 64 Carrier vehicles such as cells, liposomes, microbubbles, perfluorocarbon emulsions, and cross-linked iron oxide transport the ligands to the target molecules. In addition, carriers are also attached to signal elements that generate the signal to be detected by imaging. These include microbubbles (ultrasound), radioactive isotopes (SPECT and PET), paramagnetic and superparamagnetic compounds (MRI), iodinated compounds (CT), and fluorochromes (optical imaging).

Signal elements can be delivered in quenched form to be released only after specific enzymatic cleavage or emit a signal when the ligand binds to the target. 65 The ligand can be

Signal detector

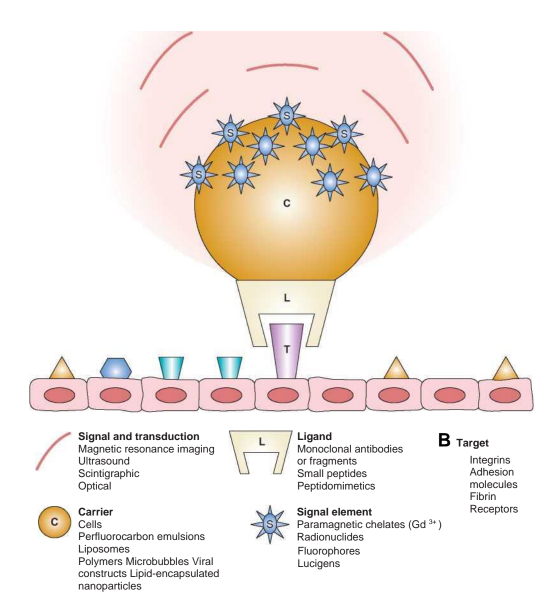


FIGURE 28-14 Schematic diagram illustrating a model of targeted contrast agents. The molecule of interest (T) is targeted by the specific ligand (L), which is conjugated to a carrier particle (C), which concentrates the signal molecules (S) at the binding site. (Modified from Choudhury RP, Fuster V, Fayad ZA: Molecular, cellular and functional imaging of atherosclerosis. Nat Rev Drug Discov 3:913, 2004.)

conveyed to the target by diffusion, mass flow, receptormediated internalization, pinocytosis, or intracellular carriage. The sensitivity of agents can be increased by the use of multivalent compounds, by conjugation of multiple ligands to a signal moiety, or by conjugation of multiple signal compounds to the target. ^{66,67}

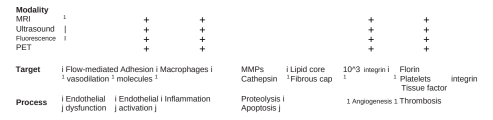
MRI has the advantages of being noninvasive and requiring no radiation, making it ideal for serial tracking. It has good spatial and temporal resolution, which is ideal for dynamic studies, and it has excellent reproducibility and is good for evaluation of multiple beds. However, it has lower sensitivity than PET or SPECT for detection of sparse targets and has prolonged imaging times.

Molecular imaging can be used at many targets (Fig. 28-15) for the diagnosis of atherosclerotic plaque, including its activity

(Table 28-6).

Molecular MRI Contrast Agents

Molecular MRI contrast agents can be broadly divided into nonspecific, targeted, and activatable types. Nonspecific agents operate by passive targeting and are used to detect physiological changes. Targeted contrast agents are used to determine structure and work by localizing proteins. Low-molecular-weight target-specific contrast agents are suitable for imaging of dense or common epitopes. Activatable contrast agents are used to localize enzymes and to determine



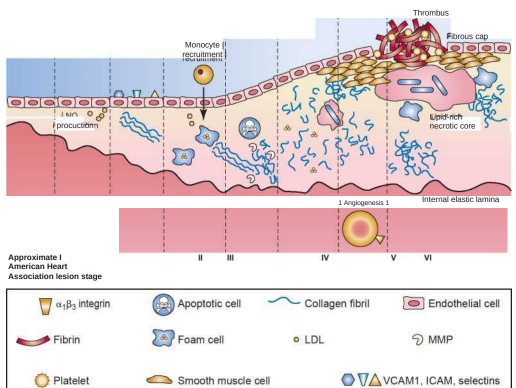


FIGURE 28-15 The various stages of atherosclerosis and the various potential targets for molecular imaging. (Modified from Choudhury RP, Fuster V Fayad ZA: Molecular, cellular and functional imaging of atherosclerosis. Nat Rev Drug Discov 3:913, 2004.)

function. Signal can be amplified and background signal minimized by use of smart agents (protease activatable, oligo-merization of signal-producing substrates), cellular trapping of phosphorylated substrates, or covalent binding. ⁶⁸

T1 Contrast Agents (Positive Agents). T1 agents, such as gadolinium, iron, and manganese, have unpaired electrons in their outer shells, which because of their paramagnetic effect produces T1 shortening of adjacent tissue protons, resulting in high signal. These agents are usually chelated to avoid the toxicity of free ions. The commonly used gadolinium agents include micelles (10 to 20 nm), lipoproteins (10 to 20 nm), and perfluorocarbon nanoparticles (200 nm). Signal from the contrast agent can be further augmented by increasing either the relaxivity or the concentration of the contrast agent.

T2/T2* Contrast Agents (Negative Agents). T2 agents produce a large dipolar magnetic field gradient because of their strong magnetic moment that accelerates dephasing of adjacent protons. The most commonly used iron-based agents have microcrystalline magnetite or maghemite cores with dextran or siloxane coating that binds to scavenger receptors in macrophages and is phagocytosed, depending on macro phage activation status. Iron-based contrast agents have both T1 and T2 shortening effects, with the predominant effect determined by the relaxivity ratio (r2/r1). If the ratio is low (ie, in dispersed agents, low concentration, or short TE/TR sequences), they produce more T1 shortening with positive contrast; but if the

ratio is high (clustered together, long TE/ TR sequences), they produce more T2 and T2* shortening with negative contrast. In addition, the extent of darkening does not directly correlate with the concentration of contrast agent because it is associated with blooming and technical parameters. No toxicity has been demonstrated so far with iron agents, and iron uptake is not inhibited by statins. ⁶⁹T2 agents are classified on the basis of their size.

- 1. Nanoparticles (10 nm-1 p m) have high relaxivity, with predominantly intravascular distribution, slow extracellular distribution, and hepatic clearance (see section later).
- 2. MPIO (microparticles of iron oxide) are the largest iron particles (0.9-4.5 p m), with high iron payload and high specificity due to less extravasation and nonspecific endothelial uptake. They distribute only in the intravascular space without any plaque uptake, making them useful for imaging of adhesion molecules ⁷⁰ and activated platelets.
- 3. SPIO (superparamagnetic iron oxide, 50-300 nm) is made up of an iron oxide core surrounded by a thin and incomplete dextran coating (eg, ferrum oxide, ferucarbotran [Resovist]). These particles usually aggregate in solution into large clusters and are rapidly cleared by reticuloen dothelial cells (t ^, 5 minutes), which limits passive diffusion, with rapid extracellular space distribution, fast renal clearance, and moderate relaxivity. Uptake in macrophages of atherosclerotic plaque has been demonstrated in mice

Processor	Target	Contrast Agent	Mechanism of Action
Endothelial dysfunction	HDL Oxidized LDL MPO	HDL-like nanoparticles (37pA,18aA) Micelles with IK17 Fab Oligomerized MPO substrates	Reverse cholesterol transport Binds to oxidized LDL Binds to MPO
Endothelial activation	VCAM-1	VINP-28 VNP Trimodality nanoparticles	Ligand homologous to VLA-4 VLA-4 homologue for MRI and NIRF MRI, NIRF, PET
	VCAM-1, P-selectin	Dual ligand	Binds to VCAM-1 and P-selectin
nflammation	Macrophages	USPIO	Phagocytosed by macrophages, T2* shortening
		Trimodality nanoparticles	Positive markers available Macrophage phagocytosis
	Scavenger receptors SRA, CD36	Gadolinium immunomicelles with antibodies to MSR	MRI, PET, NIRF Binds to MSR
Proteolysis	Matrix metalloproteinases (1, 2, 3, 8, 9, 13)	P947	Contains matrix metalloproteinase inhibitory peptides
Apoptosis	. ,	Annexin V-cross-linked iron oxide-Cy5.5	High affinity for phosphatidylserine
	Cell membrane phosphatidylserine Enzymes, cathepsins (K, B), caspases, scramblases		
Proliferation	Smooth muscle cells		
Extracellular matrix formation	Collagen, elastin, tenascin-C	Gadofluorine Elastin-binding Gd-DTPA	Binds to extracellular matrix components, elastin
Lipid core, fibrous core formation	Lipid core, fibrous cap		LDL pathway
Angiogenesis	& vp 3 integrin Increased vascularity and leakage	Paramagnetic nanoparticles Fumagillin nanoparticles Perfusion markers Gadofluorine micelles Gadofosveset	Target 0 ^A 3 integrin Theranostic agent against angiogenesis Dynamic contrast-enhanced MRI; increased enhancement
Thrombus	Fibrin Thrombus	Fibrin-binding nanoparticles EP-2104R	Peptide with high affinity for fibrin Peptide with high affinity for integrin; integrated into thrombi
	Platelets		og.atod into thombi
	Tissue factor	Peptides Antibodies to glycoprotein IIb/IIIa	
Cell trafficking		Labeled antibodies	
	Monocytes Lymphocytes Stem cells Oxidized LDL		

- and humans, 71,72 further accelerated by injection of -inflammatory cytokines. 71
- 4. USPIO (ultrasmall paramagnetic iron oxide, 20-40 nm) particles have a thick and complete dextran coating sur rounding a small nucleus of iron oxide to minimize aggregation (eg, ferumoxtran). They have a long half-life in blood (24 to 36 hours) because of small size, dextran coating, and slow reticuloendothelial system clearance, which results in passive extravasation, nonspecific uptake, and high background contrast. Uptake in atherosclerotic plaque 73 could be by passive diffusion across a leaky inflamed endothelium or endocytosis by activated blood monocytes followed by transcytosis through endothelium or transport through neovasculature. Uptake in macro phages is by pinocytosis for small particles and receptor-mediated endocytosis and phagocytosis for large particles, both of which are dependent on scavenger receptor SR-A and cytokine activation including interferon- y and interleukin -4. The intensity of ferumoxtran phagocytosis, rather than macrophage density, is decreased after treatment with p38 MAPK inhibitor. 74
- CLIO (cross-linked iron oxides, 30-50 nm) have functionalized exteriors that allow covalent conjugation of surface ligands, such as antibodies and peptides (eg, VINP-29 for VCAM-1).
- 6. VSOP (very small iron oxide particles, 4-8 nm) have a citrate coating, with electrostatic stabilization (eg, VSOP-C184).

Chemical Exchange Saturation Agents. Chemical exchange saturation agents are based on the principle of magnetization transfer. One or more pools of exchangeable protons with sharply defined resonance frequency and large chemical shifts are well separated from free water peak. Selective radiofrequency radiation of exchangeable protons results in transfer of saturated magnetization to water resonance peak and causes drop in signal intensity. These agents include small diamagnetic compounds, such as sugars and amino acids; paramagnetic lanthanides, such as europium, dysprosium, holmium, erbium, thulium, and ytterbium che lates; and macromolecular systems, such as dendrimers, polymers, single-stranded RNA, and LIPOCEST (liposomes with 0.1 mm concentration of Tm-DOTMA). 68

Fluorine 19-Based Agents. The nucleus of fluorine 19 has a high magnetogyric ratio; it is abundant in nature although minimally seen in the human body, which results in minimal background noise during imaging. Fluorinated compounds such as perfluorocarbon and perfluoropolyether oils stabilized with surfactants such as water form emulsions and can be used to label cells or function as target-specific contrast agents. ^{68 19} F MRI data can be superimposed on ¹ H MRI that depicts anatomical information.

Selective Gadolinium Agents. MRI contrast agents can be engineered to conceal gadolinium from tissue water, revealed only when a specific enzyme-mediated cleavage occurs. This has been used in imaging of β - galactosidase activity as a marker of transgene expression. Similarly, a conditionally activated probe that on exposure to myeloperoxidase causes oligomerization of gadolinium-rich generating particles with higher T1 relaxivity than a monomeric source compound has also been used. 65

Nanoparticles. Nanotechnology combines materials by precisely engineering atoms and molecules to yield new molecular assemblies on the scale of individual cells, organ cells, or even smaller (5 to 500 nm), with unique chemical and biological properties that can easily integrate multiple properties (multiple contrast, diagnostic and therapeutic, multiple targeting groups), resulting in higher payloads, higher contrast, and lengthy circulation times. Nanoparticles have a core surrounded by a coating on which are attached ligands and biocompatible polymers. Sources of contrast and therapeutics are included in the core or in the coating. ⁷⁵ The various types of nanoparticles are liposomes with lipid bilayer (50 to 700 nm); emulsions – oil in water type mixtures , stabilized with surfactants (eg, perfluorocarbons, 200 to 400 nm); polymers - flexible designer approach with controlled size (40 to 200 nm) and shape (eg, polyhydroxy acids, dendrimers); and metallic (15 to 60 nm; eg, iron oxide, gold nanoparticles, carbon nanotubes, fullerenes). However, toxicity information on nanoparticles is limited. Oxidative damage, dermal toxicity, and granulomas have been reported. 76

Theranostic Agents. Theranostic agents have integrated therapeutic and diagnostic molecules. Addition of a diagnostic moiety enables temporal and spatial monitoring of the therapeutic agent, confirms delivery at the desired target, identifies the need for dose modification, and quantifies the molecular efficiency. Nanoparticles can deliver drugs by liposomes through endocytosis or contact-facilitated drug delivery that requires close apposition between the carrier and the targeted cell membrane. Examples include fumagillin nanoparticles against angiogenesis and CLIO-THPC and CLIO-Cy5.5 against inflammation. 66 Iron oxide nanoparticles with attached Cy5.5 (near-infrared fluorescence [NIRF]), MPAP (membrane translocation myristoylated polyarginine peptide), and siRNA can be detected by both MRI and fluorescence and are efficiently taken up by cells and silence specific genes such as green fluorescence protein. siRNA attached to anti sense to survivin, an inhibitor of apoptosis proteins, is used in mice for the treatment of cancer. 68

Targets and Agents in Atherosclerosis

The various techniques of molecular MRI are described in Table 28-7.

Endothelial Adhesion Molecules. Endothelial adhesion molecules are involved in early stages of atherogenesis with increased expression. Vascular cell adhesion molecule 1 (VCAM-1) is a ligand for very late antigen 4 (VLA-4) and a $_4\,\mbox{\sc s}_1$ integrin. VINP-28 (VCAM-1 internalizing nanoparticle 28), a magnetofluorescent nanoparticle with iron oxide and the ligand VHPKQHR that is homologous to VLA-4, is trapped in VCAM1-expressing cells. Uptake in atherosclerotic plaques has been shown in mice and human carotid artery plaques. 77

Technique	Mechanism
Dynamic contrast-enhanced MRI	Assessment of permeability of neovasculature due to angiogenesis
Enhanced permeability and retention	
	Long-circulating macromolecular agent that accumulates in inflamed plaque over time due to enhanced permeability
Ex vivo labeled cells	Cells are labeled ex vivo (eg, monocytes) and injected into the body and tracked in vivo
In vivo labeled cells	
	Intravenous injection of probe or contrast agent that is taken up by cells of interest (eg, monocyte tracking)
Conjugate imaging probes with one or several targeting ligands that will recognize and bind receptors	Targeting of cell surface receptors expressed in endothelium or inside plaque with increased permeability
Modified lipoproteins	Enriched lipoproteins follow the HDL pathway and localize to the lipid core

Another ligand, VHHSPNKK, with homology to a chain of VLA-4, also showed similar results. ⁷⁸ VINP-28 can also be conjugated to fluorescence, infrared, or radioactive probes. The trimodality nanoparticle for detection by MRI, PET, and NIRF is made up of dextrinated and DTPA-modified nanoparticles (20 nm), labeled ⁶⁴ Cu (PET tracer), and a fluorophore. ⁷⁹ Dual ligand technique (binding to VCAM-1 and P-selectin) has also been demonstrated.

Inflammation. Inflammation plays an important role in the pathogenesis of atherosclerosis, resulting in plaque rupture, thrombotic vessel occlusion, and major vascular events. Macrophages are involved in atheroma initiation, propagation, and rupture.

USPIO or magnetic nanoparticles coated with dextran, such ferumoxtran-10 and ferumoxytol, are taken up by macrophages in atherosclerotic plaques and produce dark signal in MRI due to T2* shortening, 69 with good histopatho logical correlation to plaque macrophage activity and iron deposition in carotid endarterectomy specimens 81 but no correlation with luminal stenosis, indicating that they are independent risk factors. In stroke patients, USPIO uptake was seen even on the asymptomatic side; and in patients undergoing coronary artery bypass grafting, uptake was seen in asymptomatic atheromas, indicating the systemic nature of the disease. ^{69,81} Vessel inflammation may be quantified to identify high-risk plaques. Imaging of coronary plaques by this technique would require high resolution and signal-to-noise ratio, probably achievable with intravascular coils or novel pulse sequences. ⁷⁰ Bimodality nanoparticles (CLIO-Cy5.5, CLÎO-VT680, CLIO-Cy7, CLIO-VT570) for NIRF and MRI and trimodality nanoparticles (64 Cu, CLIO-VT680) for NIRF, PET, and MRI are available and have higher signal and sensitivity. 79

USPIOs have been used in plaque progression and regression trials, both in animals and in humans. Anti-inflammatory - treatment reduced USPIO uptake by 70% in a mouse model. In the ATHEROMA study, in which subjects received either 10 mg or 80 mg of atorvastatin, imaging at 12 weeks with ferumoxtran showed significant reduction of inflammation in the higher dose group (Fig. 28-16). 82 However, these studies are based on visual estimation, and quantitative techniques are required. In another study, 74 an anti-inflammatory agent, p38 MAPK (mitogenactivated protein kinase) inhibitor SB-239063, was given to knockout mice, which resulted

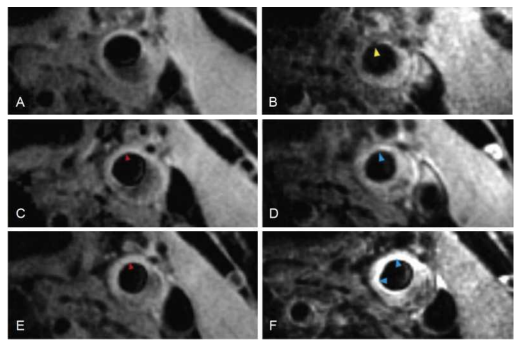


FIGURE 28-16 T2*-weighted imaging of a left common carotid artery before and after ultrasmall superparamagnetic iron oxide (USPIO) infusion at the three time points of 0 (A and B), 6 (C and D), and 12 weeks (E and F). B, USPIO uptake can clearly be seen in the plaque at baseline (arrowhead). C and E, Pre-USPIO imaging remains similar at all three time points with Sinerem having been cycled out of the plaque before reimaging (arrowhead). D, The plaque begins to enhance at 6 weeks (arrowhead). This signifies that there is a predominant T1 effect, indicating minimal USPIO uptake and a lack of activated macrophages (minimal inflammation). E, No residual USPIO signal is also seen in the pre-USPIO imaging at 12 weeks. F, Signal enhancement post-USPIO can be seen with no evidence of signal voids (arrowheads). (Modified from Tang TY, Howarth SPS, Miller SR, et al: The ATHEROMA [Atorvastatin Therapy: Effects on Reduction of Macrophage Activity] Study. Evaluation using ultrasmall superparamagnetic iron oxide-enhanced magnetic resonance imaging in carotid disease. J Am Coll Cardiol 53:2039, 2009.)

in no change in plaque macrophage content between SB-239063 animals and normal controls, but diminished iron oxide uptake on MRI was seen in SB-239063-treated mice, suggesting a decrease in macrophage phagocytic activity. ^{69.81}

The signal produced by iron agents is highly dependent on technical parameters. In addition, signal is limited by long delay for imaging and background imaging artifacts. Bright signal can be obtained with the use of iron agents, in appropriate sequences and concentration. Serial inversion recovery MRI with monocrystalline iron oxide nanoparticle-47 produced high signal in rabbits. 83 Other techniques include gradient echo acquisition for SPIO, inversion recovery on resonance water suppression, and ultrashort echo time, all of which enhance the signal. 84 Postcontrast T1-weighted MRA done 5 days after introduction of USPIO will generate high signal in the lumen because of low concentration of luminal USPIO (T1 shortening) at 5 days and signal drop in the plaques because of high concentration in plaque (T2/T2* shortening), 73 which is ideal for visualization of both the lumen and the plaques. Another interesting study found that in carotid stenosis, dark signal is seen in symptomatic patients because of high macrophage content, high uptake, and high concentration, but mild high signal is seen in asymptomatic patients because of large fibrous caps, fewer macrophages, and low concentration of USPIO. 85

Nanoparticles containing immunomicelles with monoclonal antibodies targeting scavenger receptors (MSR) and high payload of gadolinium (5900 molecules) have been used in mice (Fig. 28-17) ⁸⁶ to demonstrate high signal at 1 and 24 hours in actively inflamed plaques, with good histopatho logical correlation. USPIOs can potentially be used in the future to target macrophage markers such as MAC387 and CD68. ⁶⁹

Angiogenesis. Angiogenesis results in enhanced -permeability of neovasculature, which can be detected by dynamic contrast-enhanced MRI (DCE-MRI) or by use of long-circulating contrast agents that progressively accumulate in the plaque.

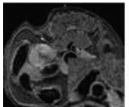
Dynamic Contrast-Enhanced MRI. Post-gadolinium contrast enhancement kinetics measured by DCE-MRI has been shown to correlate with plaque neovascularity and vulnerability . Kinetic modeling and integrated area under the enhancement versus time curve are useful in characterizing enhancement dynamics in plaque. Fractional plasma volume (V p) correlates strongly with histologically validated neovas culture content, which is the main source of entry of inflammatory cells into atherosclerotic plaques. Transfer constant (R transs) is also a potential marker of plaque inflammation because it correlates with macrophage density, ⁸⁷ neovasculature, ⁸⁸ loose matrix content, serum markers of inflammation, and proinflammatory risk factors (C-reactive protein, LDL, smoking). There is mild overestimation of neovasculature as measured by dynamic MRI because of rapid exchange of contrast agent between the plasma and interstitial volumes. Any region that comes to equilibrium within one time frame of dynamic sequence will be indistinguishable from blood 13 (Fig. 28-18).

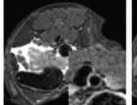
Macromolecular Agents. Macromolecular agents such as gadofluorine and paramagnetic micellar agents accumulate over time in atherosclerotic plaques because of increased vascular permeability and retention. In addition, because of its lipophilic nature, gadofluorine forms 5-nm micelles in aqueous solution and preferentially labels the fatty cores of plaques in cholesterol-fed rabbits. ⁸⁹

Gadofosveset (Vasovist, MS-325) is a new gadolinium-based contrast agent that heavily (> 75%) binds to serum

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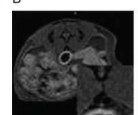


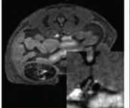


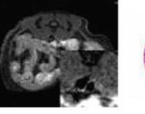




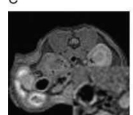


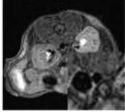












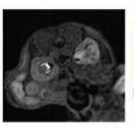




FIGURE 28-17 In vivo MR images obtained at baseline and after injection of macrophage-targeted immunomicelles (A, B), untargeted micelles (C), and Gd-DTPA (D) in ApoE-mice. The MRI insets are enlargements of the aortas.

A-D, right, Hematoxylin and eosin sections of the aorta at the identical anatomic level as the MRI images from the same animal. Very significant heterogeneous enhancement of the aortic wall was seen, at least twofold higher than untargeted micelles. (Modified from Amirbekian V, Lipinski MJ, Briley-Saebo KC, et al: Detecting and assessing macrophages in vivo to evaluate atherosclerosis noninvasively using molecular MRI. Proc Natl Acad Sci U S A 104:961, 2007.)

albumin (> 75%), resulting in a long elimination half-life and high relaxivity. It produces higher enhancement than gadopentetate dimeglumine (Gd-DTPA) in atherosclerotic plaques of rabbits, either because of increased entry into a leaky microvasculature due to its small size or through normal microvasculature due to albumin binding (Fig. 28-19). Once inside the plaque, it binds to albumin, which is present in higher quantities in plaques, resulting in increased relaxivity, which causes increased signal. ⁹⁰

Gadofluorine M (GdF) enhancement correlates directly with plaque instability because it preferentially binds to collagenous material within plaques, and it accumulates more and penetrates deeper into plaques with large numbers of foam cells. ⁹¹ It also has a long plaque half-life (24 hours), which improves the conspicuity of atherosclerotic vessel wall both at 1 and 24 hours. Multiple plaques and multiple vascular territories can be imaged with a single injection.

Angiogenesis Marker Imaging. a $_{\rm v}$ p $_{\rm 3}$ integrin (glyco-protein)-specific antibodies and a $_{\rm v}$ p $_{\rm 3}$ -specific RGD peptides or peptidomimetics are molecules that target markers expressed on angiogenically activated endothelial cells. Nanoparticle targeted to a $_{\rm v}$ p $_{\rm 3}$ integrin is a perfluorocarbon type, with a high payload of gadolinium (Gd-DTPA bis-oleate, 90,000 atoms per particle) that can be imaged by both ultra sound and MRI. 68

Theranostic Agent. Fumagillin, an antiangiogenic drug, has been delivered along with integrin-targeted nanoparticles in rabbits, which showed decreased MRI enhancement 1 week after treatment. Baseline enhancement predicted therapeutic response, with low response in those with higher base line enhancement. Dual angiogenesis-targeted fumagillin nanoparticles (integrin a $_{\rm V}$ p $_{\rm 3}$, a $_{\rm 5}$ p $_{\rm i}$) is more effective than single. Fumagillin has also been shown to have a synergistic antiangiogenic effect with statins. 92

Proteases. Inflammatory proteases promote extracellular matrix digestion, plaque remodeling, and fibrous cap rupture.

Matrix Metalloproteinases. Matrix metalloproteinase is a surrogate marker for the presence of macrophages. P947 is a gadolinium chelate (Gd-DOTA) covalently bound to a peptide that specifically binds to matrix metalloproteinases at the

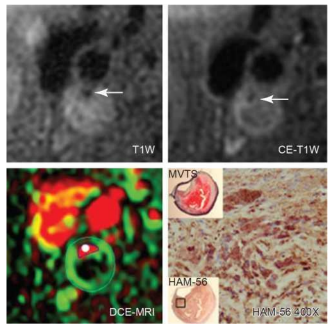


FIGURE 28-18 Dynamic contrast-enhanced MRI of an inflamed fibrous cap of plaque (in T1W and CE-T1W) shows high transfer constant (green) in the adventitial vasa vasorum (near the outer vessel wall boundary) and along the lumen boundary (inner boundary demarcating red lumen region). (Modified from Chu B, Ferguson MS, Chen H, et al: Cardiac magnetic resonance features of the disruption-prone and the disrupted carotid plaque. JACC Cardiovasc Imaging 2:883, 2009.)

enzymatic active site; it has been shown to localize to the fibrous cap of atherosclerotic plaque in mice and human carotid arteries, producing greater and longer contrast effects than Gd-DTPA. ⁸⁷, ⁹³

Myeloperoxidase. Myeloperoxidase is a heme peroxidase enzyme that generates oxidant species hypochlorous acid. It is produced by macrophages within the plaque and promotes atherogenesis by modifying LDL, inactivating HDL and nitric oxide, activating matrix metalloproteinases, releasing tissue factor, and causing apoptosis. Myeloperoxidase substrates are available that, when oligomerized, produce high signal with MRI. ⁶⁶

Cathepsins. Cathepsins include cysteine proteases that are produced by macrophages, endothelial cells, and smooth muscle cells. Cathepsins S and K degrade extracellular matrix through their elastase and collagenase properties and destabilize plaque. Cathepsin K localizes to the shoulder of ruptured atherosclerotic plaques, which can be imaged by NIRF or MRI with an activatable imaging agent that becomes fluorescent after enzymatic cleavage by cathepsin B and is detected by fluorescence-mediated tomography co-registered with MRI. 94

Apoptosis. Apoptotic cells express on their surface phosphatidylserine, a phospholipid that is normally seen in the inner cell membrane of viable cells. Annexin A binds to phosphatidylserine, which, when it is cross-linked to iron oxide, can be used for MRI in humans and animals. Dual modality particles (CLIO-annexin-Cy5.5) use iron oxide and Cy5.5 for MRI and optical imaging. ⁶⁸ Annexin V has been shown to localize with nonapoptotic macrophages and intra plaque hemorrhages.

Extracellular Matrix Production. Elastin-binding Gd-DTPA, a novel low-molecular-weight gadolinium chelate (¹⁵³ Gd-labeled BMS753951) with high affinity to elastin and a high relaxivity, localizes preferentially in the arterial wall of hypercholesterolemic rabbits, facilitating MRI imaging of plaque location, lesion burden, and remodeling. Uptake was seen in ApoE mice at the sites of extracellular matrix

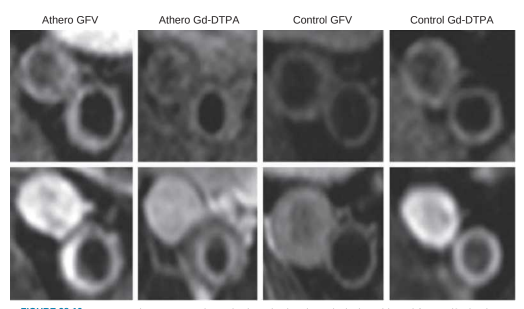
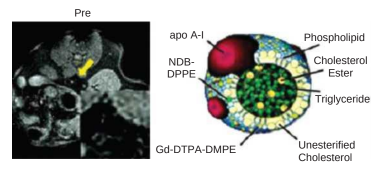


FIGURE 28-19 Precontrast and postcontrast MR images in atherosclerotic and control animals receiving gadofosveset (GFV) and Gd-DTPA. Top row, Precontrast images. Bottom row, Postcontrast images. Athero, atherosclerotic rabbit; Gd-DTPA, gadopentetate dimeglumine. Signal enhancement after contrast agent administration is seen in all groups except control animals receiving gadofosveset. The signal enhancement is higher for atherosclerotic than for control animals imaged with gadofosveset. Gd-DTPA could not enable discrimination between normal and atherosclerotic vessel walls. (Modified from Lobbes MBI, Miserus RJ, Heeneman S, et al: Atherosclerosis: contrast-enhanced MR imaging of vessel wall in rabbit model—comparison of gadofosveset and gadopentetate dimeglumine. Radiology 250:682, 2009.)





FIGURE 28-20 Sagittal black blood MRI (A) after injection of EP-2104R shows high uptake within thrombus (arrow) in descending thoracic aorta. In the CT scan (B), plaque is seen as a dark structure (arrow). (Modified from Spuentrup E, Botnar RM, Wiethoff AJ, et al: MR imaging of thrombi using EP-2104R, a fibrin specific contrast agent: initial results in patients, Eur Radiol 18:1995, 2008.)



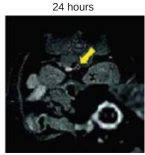


FIGURE 28-21 In vivo MR images of abdominal aorta in 8-week-old mouse, before (left image) and after (right image) injection of recombinant HDL-like nanoparticles (middle image), show uptake within the abdominal aortic plaque (arrow). (Modified from Frias JC, Williams KJ, Fisher EA, et al: Recombinant HDL-like nanoparticles: a specific agent for MRI of atherosclerotic plaques. J Am Chem Soc 126:16316, 2004.)

formation (intimal hyperplasia), aneurysm (elastin degradation), and hypertension (excess elastin formation), including coronary artery plaques and post-stent injuries. ⁹⁵

Thrombus. Thrombus can be imaged with the use of ligands against targets such as fibrin, platelet integrins, tissue factor, and factor XIIIa.

Fibrin-specific probes using paramagnetic perfluorocarbon nanoparticles with a high payload of gadolinium chelates (50,000 to 90,000 gadolinium atoms) conjugated with fibrin specific antibodies localize to disrupted carotid plaques. ⁹⁶ Fibrin-binding gadolinium-labeled peptide with only four gadolinium chelates per targeting moiety can also be used if there is abundant fibrin. Iron particles with antifibrin antibodies have also been used in animal studies. ⁹⁷

EP-2104R is a novel fibrin-specific gadolinium-based agent that localizes to and enhances in the intraluminal thrombus layer. With use of this agent, a vulnerable plaque with plaque erosion and thrombus can be localized with single delayed postcontrast heavily T1-weighted electrocardiography-gated black blood sequence ⁹⁷ (Fig. 28-20). Thrombi of various ages can be localized in various vascular territories and the heart. In humans, its use has been demonstrated in patients with transient ischemic attack, but coronary arterial imaging would be challenging.

Platelets could be targeted by arginine-glycine-aspartic acid peptides or single-chain antibody that is specific for activated conformation of glycoprotein IIb/IIIa. Tissue factor and factor XIIIa have also been imaged in vitro with MRI. ⁷⁶

Lipoproteins. Enriched lipoproteins with contrast-inducing species can be used to localize to the lipid core because they will follow the HDL pathways into plaque macrophages. HDL for molecular imaging can be reconstituted from humans or can be recombinant. Reconstituted HDL particle (9 nm) contains

phospholipids with human apolipoprotein AI, with or without unesterified cholesterol and Gd-DTPA-DMPE. Recombinant HDL is made up of apolipoprotein AI-mimicking peptide 37pA (7.6 nm) or 18aA (8.0 nm), phospholipid, Gd-DTPA-DMPE (15 to 20 molecules per particle), and fluorescent phopholipid ⁹⁸ (Fig. 28-21). Both these molecules are localized to the intima of atherosclerotic plaques in mice and enhanced in 24 hours. ⁹⁹

Oxidized LDL is a key factor in the initiation and progression of atherosclerosis that indicates an unstable plaque because it upregulates matrix metalloproteinases and apoptosis. Micelles containing gadolinium and IK17 Fab (antibodies to oxidation-specific isotopes) localize to atherosclerotic plaques in apolipoprotein E-deficient mice at 72 hours. ¹⁰⁰

Osteogenesis and Calcification. The superficial calcified nodule is a marker of vulnerable plaque, and this can be detected by macrophage-avid MRI magnetofluorescent nanoparticles coupled with intravital confocal fluorescence microscopy. Bisphosphonate probes have been used to detect osteogenesis in early-stage carotid plaques. ¹⁰¹

POSITRON EMISSION TOMOGRAPHY

PET is based on beta decay of radioisotopes that results in the emission of a positron, a positively charged beta particle. After emission, the positron travels a few millimeters in tissue and collides with an electron, which results in complete annihilation of both the positron and electron and conversion to energy in the form of two high-energy (511 keV) gamma rays that are released at 180 degrees from each other.

PET detectors register only events with temporal coincidence of photons striking at directly opposite detectors, which enhances the spatial and temporal resolution of PET compared with SPECT. Easy labeling of primary substrates for energy metabolism and membrane receptor subtypes in the heart allows highly sensitive investigation of physiological pathways and quantization of tissue metabolism.

PET uses positron-emitting isotopes (oxygen 15, carbon 11, nitrogen 13, and fluorine 18) that are incorporated into physiologically active molecules. Rubidium Rb 82 chloride, [13 N]ammonia, and [15 O]water are tracers that evaluate perfusin* [18 F]flnorodeowcrliicose (FDG) [11 C]acetate ad [11 C] sion, [r]uoioeoxygiucose (FDG), [C]aceiaie, an [C] palmitate evaluated metabolism. Dynamic-mode PET allows evaluation of rate of change of physiological processes. However, PET scanning is associated with significant ionizing radiation and has limited

spatial resolution to evaluate coronary arteries.

PET/CT is a type of hybrid imaging that combines the images from PET and CT scans performed at the same time on the same machine, which helps in correlation of anatomical and functional information, thereby improving accuracy, at the same time, it reduces scanning time (faster CT acquisition for co-registration and attenuation correction of PET data) and radiation dose. Multiple targets can also be evaluated in the same examination. ³⁹ PET/MRI co-registers low spatial resolution PET with high spatial resolution MRI images, with significantly lower radiation dose than in PET/CT.

In the cardiovascular system, PET can be used in direct evaluation of the atherosclerotic plaque and also in the assessment of myocardial ischemia.

PET Imaging of Atherosclerotic Plaque

Imaging of Macrophages

FDG is taken up into cells similar to glucose, but it does not have the metabolic pathway for further metabolism and excretion. Linked to ¹⁸ F, a positron emitter, FDG can be used to detect and quantify FDG within the tissue of interest. FDG-PET is useful in assessment of inflammatory activity within atherosclerotic plaque because the macrophages in inflamed plaque show avid FDG uptake and accumulation. The efficacy of PET in detecting atherosclerotic inflammation in various vascular beds has been extensively demonstrated in various human and animal 5 studies that used PET/CT or co-registered information with CT or MRI.

Higher FDG uptake has been demonstrated in symptom atic than in asymptomatic carotid plaques. 102 Autoradiogra phy confirmed that the uptake was within macrophages, and there was a direct correlation between FDG uptake and macrophage burden in the plaque (Fig. 28-22). 103 The carotid FDG uptake was higher in men, older age, and those with higher inflammatory biomarkers, metabolic syndrome, hypertension, hyperlipidemia, low HDL concentration, high levels of high-sensitivity C-reactive protein, and CAD. 104,105 In symptomatic patients with transient ischemic attack, uptake was also seen in asymptomatic nonstenotic areas on the opposite and same side, indicating that angiography may not always identify the culprit lesion. ¹⁰⁶ There was no overlap between inflammation and calcification, indicating that calcification represents a much later stage of the disease process.

Combined MRI and PET can identify lesions responsible for embolic events, with MRI delineating the vascular lumen and PET identifying inflammation. 107 In addition, FDG uptake may predict plaque rupture and clinical events. In a rabbit atherosclerotic model, plaque thrombosis was promoted by Russell viper venom injection, and FDG-PET was done before and after triggering of thrombus. Segments that developed thrombus had the highest FDG uptake and highest macrophage activity. FDG uptake and hence plaque inflammation

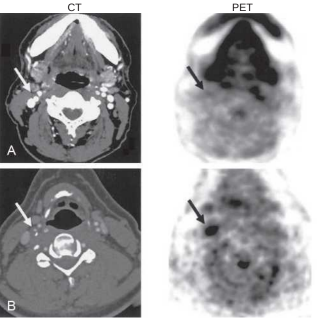
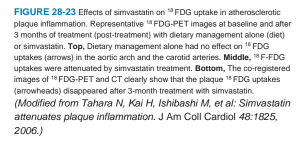


FIGURE 28-22 Axial PET images and the co-registered CT images from two patients, one (patient A) who manifested low ¹⁸F-fluorodeoxyglucose (FDG) uptake in the region of the carotid plaque and one (patient B) with high FDG uptake in the region of the carotid plaque. The region of the excised carotid plaque is noted with arrows. A, Trichrome-stained histologic specimen from patient A demonstrated a collagen-rich plaque with low lipid content, and CD68 staining on the high-powered images demonstrated limited macrophage infiltration. These histologic features are consistent with a metabolically stable and potentially clinically stable plaque. B, Trichrome-stained histologic specimen from patient B demonstrated a complex plaque with a necrotic core, and the CD68 staining demonstrated intense macrophage infiltration. These histologic features are consistent with a metabolically unstable plaque that is vulnerable to rupture. (Modified from Tawakol A, Migrino RQ, Bashian GG, et al: In vivo 18Ffiuorodeoxyglucose positron emission tomography imaging provides a noninvasive measure of carotid plaque inflammation in patients. J Am Coll Cardiol 48:1818, 2006.)

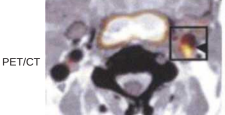
In a study of 2000 cancer patients, patients with the highest $\frac{0}{2}$ arterial FDG uptake were more likely to have previously suffered m a vascular event. 109

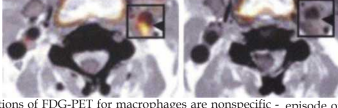
FDG-PET is a proven, reproducible technique in the detection of inflammation in different arterial beds, such as the aorta 110 and the vertebral, 107 brachial, subclavian, and peripheral arteries, including femoral and iliac arteries. 111 However, PET imaging of coronary arteries is limited by significant background FDG uptake by metabolically active myocar dium, small size of the coronary arteries and plaque, 112 and deep location, tortuosity, and mobility of coronary arteries. Myocardial FDG uptake could be bypassed by use of high-fat/low-glucose protein preferred diet or beta blockers before imaging. Spatial resolution can be improved by co-registering with CT or MRI or by using intravascular radiation detectors. Electrocardiographic or respiratory gating is used to reduce motion. 110

PET has also been well established as a useful technique in serial follow-up for evaluation of plaque for progression or regression, with high interobserver and intraobserver reproducibility, 5 although there might be a change in pattern of uptake due to waxing and waning of atherosclerosis. 113 Decreased FDG uptake has been demonstrated in the aortic wall of atherosclerotic rabbits after the use of statins 114 and after 3month treatment with probucol. 115 In humans, reduction of carotid plaque inflammation that correlated with HDL elevation was demonstrated after just 3 months of simvastatin therapy in 43 oncology patients (Fig. 28-23). MRI typically takes up to a year for changes to be visualized. 116









artifact. This could be overcome by concomitant use of high-solely on symptoms and morphological detection of stenosis. resolution MRI/CT.

Other Macrophage Targets

Other targets for imaging of macrophages include benzodiazepine in mitochondrial membranes, macrophages, using targeted agents such as [11 C](R)-PK11195 117 moderate exercise and stress management reduced size and particularly or PBR28. 70

Other PET Targets

Proliferating smooth muscle cells in the plaque can be imaged by indium In 111-labeled Z2D3. LDL labeled with indium In 125 and epicardial coronary arteries decrease downstream blood flow and technetium Tc 99m accumulates in the lipid core of atherosclerosis perfusion, resulting in small areas of subendocardial necrosis, and can be imaged. Annexin V and other markers can also be used, which eventually leads to fibrosis and left ventricular dysfunction. labeled with PET-compatible tracers such as ¹²⁴ I and ¹⁸ F.

VCAM

With use of ¹⁸ F-labeled 4V (linear, multivalent tetrameric synthetic peptide VHPKQHR, which has sequence homology to VLA-4 and increased affinity to VCAM-1), PET/CT showed high uptake in endothelium of atherosclerotic lesions in hypercholesterolemic mice and low uptake in mice receiving an atorvastatin-enriched diet. 118 This tracer has good potential for human use because of its short circulating half-life and high atherosclerotic/nonatherosclerotic lesion uptake.

RAGE (receptor of advanced glycation end products) is an important target in the vessel wall that has been implicated in atherosclerotic progression. It belongs to the immunoglobulin superfamily of cell surface receptors that is expressed in endothelial cells, smooth muscle cells, monocytes-macrophages, and lymphocytes. The three important RAGE ligands are advanced glycation end products, S100/calgranulins, and high-mobility group box 1/amphoterin. The interaction of RAGE ligands with RAGE in inflammatory cells leads to activation of transcription factors, which leads to upregulation of cytokines, adhesion molecules, matrix metalloproteinases, and tissue factor, resulting in chronic cellular activation, vascular inflammation, and endothelial dysfunction. 99m Tc-labeled anti-RAGE F(ab ') 2 antibodies have been used in atherosclerotic mice. 119 PET studies have been done in rats with use of recombinant human S100A1, S100B, and S100A12 proteins labeled with the positron emitter fluorine 18 (18 F) by conjugation with A -succinimidyl-[¹⁸ F]fluorobenzoate ([¹⁸ FJSFB). 120

of Risk Factor Reduction

The majority of the burden of CAD is accounted for by traditional risk factors. 121 Asymptomatic silent ischemia is the most common manifestation of CAD and a predictor of adverse clinical outcomes, such as coronary events and cardiac death. 122 Silent ischemia carries a similar adverse prognosis in patients after an

The limitations of FDG-PET for macrophages are nonspecific - episode of unstable angina, 123 MI, 124 or chronic stable angina. 125 uptake in endothelial cells and lymphocytes and partial volume Decisions for invasive workup and intervention cannot be based

> Hyperlipidemia-induced endothelial dysfunction can also cause vasoconstriction of established coronary stenosis and result in silent or overt ischemia, 126 which could be partially reversed by use of a statin. 127 Low-saturated fat, vegetarian diets with mild to severity of perfusion abnormalities on PET scan with only modest regression on angiographic score. 128 Stress reduction has also been shown to treat ischemic episodes. 129

> Repeated episodes of silent ischemia caused by obstruction of Cardiac risk increases with severity of ischemia. ¹³⁰ Functional tests are necessary to identify the presence, extent, and severity of myocardial ischemia, which is associated with outcome. A normal perfusion scan excludes myocardial ischemia, with an event rate of less than 1% per year. 131 These tests can act as a gatekeeper for coronary angiography and help in choosing the optimal therapeutic pathway.

> Ischemia can be managed by either anti-ischemic drugs (beta blockers, calcium channel blockers, nitrates, or a combination) or revascularization procedures. There is no strong evidence to suggest that complete suppression of ischemia with aggressive drug therapy improves the adverse clinical

28

ischemia after revascularization. The Asymptomatic Cardiac vasodilation). Coronary functional and microvascular alterations Ischemia Pilot (ACIP) study 132 found that coronary artery bypass may coexist with epicardial coronary artery lesions and contribute grafting suppressed ischemia more than percutaneous - to ischemia. In addition, endothelial dysfunction without transluminal coronary angioplasty at 12 weeks, and 1- or 2-year significant coronary artery stenosis can also cause ischemia by mortality was lower for revascularized patients than for those coronary vascular and blood flow abnormalities that reduce receiving medical therapy. The DANAMI 133 and SWISSI II 134 trials coronary vasodilating MH properties at macrovascular and showed better outcomes for percutaneous coronary intervention microvascular levels. In addition, a stenosis may not be discrete. compared with medical therapy for post-MI ischemia, although Development of lateral neck also changes the blood flow patterns. both trials did not use optimal medical therapy or optimized interventional therapy. The COURAGE trial compared optimal medical therapy with optimal medical therapy and percutaneous Role of PET in the Evaluation of Myocardial coronary intervention with bare metal stents and found no Ischemia difference between both groups in the primary endpoint of death from any cause and nonfatal MI in 4.6 years. 129

Investigative procedures are necessary for objective assessment of therapeutic response to ischemia. Although Holter monitoring and elimination of ST-segment depression on electrocardiography can be used to evaluate the efficacy of these therapies, they are not accurate or reliable because of marked variability in the number and duration of ischemic episodes as detected by this technique. 135 PET is an ideal technique for evaluation of silent or overt ischemia and assessment of response to various therapies because of its high accuracy and reliability in quantitative estimation of myocardial blood flow and perfusion reserve.

Detection of myocardium previously exposed to ischemia can be useful in identification of myocardium at risk for acute coronary syndromes. Annexin V and 123 I-labeled S -methyl- p -iodophenylpentadecanoic acid have potential ischemic molecular markers. 130

Evaluation of Ischemia by PET

Coronary Blood Reserve

Coronary or myocardial blood reserve (MBR) is the increase of myocardial blood flow (MBF) from rest to stress (MBR = MBF during stress - MBF during rest). Coronary blood flow normally adapts rapidly to meet the changing myocardial oxygen demands to maintain normal contraction. Because oxygen extraction at rest limitation of PET because exercise is an important component of is already at maximum, oxygen supply can be maintained at stress myocardial perfusion imaging studies with independent only by increasing coronary blood flow. Coronary blood flow prognostic and diagnostic value (Fig. 28-24). depends on aortic diastolic pressure and downstream resistance. Aortic diastolic pressure does not vary much from rest, and hence blood flow can be maintained only by decreased resistance.

In animals, resistance to coronary flow is provided by large epicardial vessels (R1), coronary arterioles (R2), and wall tension from ventricular chambers (R3). R2 predominates at rest, but with stress, blood flow increases up to four times because of reduction of R2 resistance and mild dilation of epicardial vessels with normal endothelial cell function. R3 may be increased or unchanged, depending on the increase in chamber radius and wall tension. In mild coronary stenosis, flow is maintained at rest and stress by autoregulatory dila tion of downstream arteriolar resistance vessels. With moderate stenosis, rest flow is maintained by use of coronary reserve, but inadequate vasodilatory reserve is available during stress to decrease resistance and to increase flow. With severe stenosis, there is no vasodilatory reserve to maintain even resting circulation.

In humans, CAD is more complex, with the coronary blood reserve affected by the length and complexity of the stenosis. The [18 F]FDG presence and extent of CAD are not always related to clinical FDG is used for imaging of myocardial glucose use. FDG manifestations of ischemic heart disease, and it is not always exchanges across capillaries and cell membranes and is possible to predict its progression and response to treatment. In humans with normal endothelial function, during stress, coronary flow increases because of coronary arterial and

arteriolar vasodilation, resulting in maximal coronary flow 483 reserve.

Subjects with traditional coronary risk factors demonstrated abnormal coronary vasoreactivity in both epicardial and resistance

outcomes. Several studies have shown improved outcomes in silent vessels (through passive elastic behavior of micro-vascular walls at

PET plays a vital role in the evaluation of patients with cardio vascular risk factors or cardiovascular disease and is a wellestablished modality for evaluation of myocardial perfusion, viability, and metabolism. 136 SPECT measures only regional differences in flow. Balanced ischemia due to multivessel CAD will not be detected on SPECT because of uniform decrease in flow reserve in all vascular territories. PET can quantify absolute regional coronary blood flow at rest and stress and calculate the coronary blood flow reserve.

Detection of mild abnormalities in myocardial blood flow reserve with PET potentially allows early identification of CAD that is characterized by endothelial dysfunction in asymptomatic patients with elevated cholesterol, smoking, hypertension, and insulin resistance. Abnormal blood flow reserve may be a predictor of future cardiovascular outcomes among patients with cardiomyopathies in the absence of CAD. 137

Compared with SPECT, PET has higher spatial resolution, improved attenuation and scatter correction, and higher sensitivity and specificity in detection of CAD. PET also provides information on cardiac metabolism in absolute terms. The main limitation of PET is the short half-life of the tracers, [13 N]ammonia and 82 Rb, which necessitates an on-site cyclotron for [13 N]ammonia and expensive monthly replacement of the generator for 82 Rb. Use of drugs instead of exercise is another

Cardiac Metabolism

Fasting Metabolism

ATP is generated by oxidative phosphorylation and glycolysis. The major energy sources of the heart are fatty acids, glucose, and lactate, depending on the physiological condition and arterial concentrations. During fasting, energy is mainly derived from long-chain free fatty acids and to a lesser extent (15% to 20%) from glucose. With normal oxygen supply, ATP and tissue citrate formed by breakdown of fatty acids suppress glucose oxidation. With decreased oxygen supply, ATP and citrate levels fall, and glycolysis is accelerated. Anaerobic glycolysis is maintained only if lactate and hydrogen ions are removed without accumulation. In severe hypoperfusion, lactate and hydrogen accumulate, which inhibits glycolytic enzymes, thus depleting high-energy phosphates, resulting in cell membrane disruption and death.

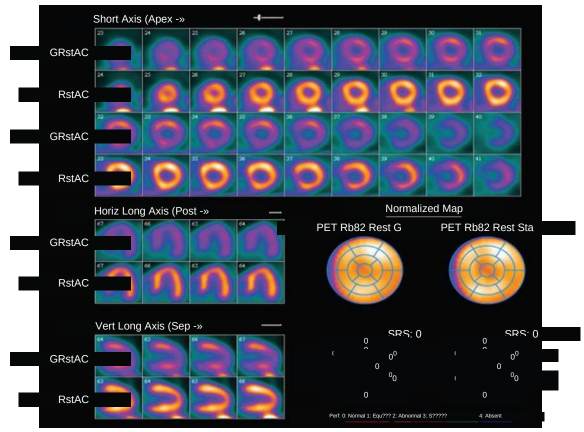


FIGURE 28-24 Normal appearances of PET perfusion scan using 82Rb in a 52-year-old man. No defect is seen in either the rest scan or the post-dipyridamole 82Rb stress scan, indicating absence of ischemia.

phosphorylated by hexokinase to FDG 6-phosphate, but it is not metabolized further or used in glycogen synthesis and is trapped higher future CHD risk, facilitated by underlying endothelial in the myocardium because of slow dephosphorylation, thus dysfunction that promotes coronary vasoconstriction and allowing imaging. PET scan is performed after loading of 50 to 75 g thrombosis. Clinically overt abnormalities in flow reserve mean of glucose 1 to 2 hours before injection of 5 to 10 mCi of FDG. In advanced coronary atherosclerosis and deserve aggressive risk diabetics, image quality may not be good because of low plasma factor modification and possible revascularization. However, insulin, noninhibition of tissue lipolysis, and high free fatty acid preclinical abnormalities in vasodilator reserve on quantitative PET levels. Intravenous insulin after glucose loading, hyperinsulinemic (but with low to intermediate risk based on Framingham score) also euglycemic clamping, and nicotinic acid derivatives are used in relate to higher estimated CHD risk and might necessitate more diabetics.

PET in Early-Stage Coronary Artery Disease

Abnormal impairment of myocardial perfusion reserve (MPR) without angiographically demonstrated coronary stenosis indicates either undetected atherosclerosis or coronary microvascular or endothelial dysfunction after hemodynamic and extravascular factors are excluded (Fig. 28-25). Myocar dial blood flow and MPR can be globally reduced in remote myocardium supplied by angiographically normal coronary arteries in patients with CAD elsewhere. Few studies ¹³⁷ have shown preserved MPR in regions supplied by angiographically normal coronary arteries in patients with one-vessel CAD. This information is useful to stratify prognosis, to determine treatment, and to monitor

MPR measured by PET is inversely related to 10-year CHD risk PET in Advanced Cardiovascular Disease in a population with no known CAD but with low to intermediate. In patients with established CAD, PET identifies patients with CAD risk based on Framingham score. ¹³⁸ This reflects preclinical disease, which could be a combination of subclinical atherosclerosis and endothelial dysfunction, either of which is not severe enough to be manifested as an overt perfusion defect (in SPECT scan) but is detected by more sensitive measure of myocardial blood flow with PET; 50% of patients with normal myocardial perfusion PET images show evidence of non-flow-limiting coronary atherosclerosis on CT scan.

Progression of subclinical disease over time contributes to aggressive risk factor modification for these subjects. MPR provides a way to document how risk factors translate into measurable damage to coronary circulation, predicting future cardiovascular events, independent of significant coronary

In patients with known CAD, there was a significantly worse prognosis in patients with higher reduction of global MPR, making it a more sensitive predictor of sudden death and other adverse outcomes than left ventricular ejection fraction. The prognosis was dependent on global impairment of myocardial perfusion and independent of the extent of regional ischemia. This global impairment is common in left ventricular dysfunction due to either primitive myocardial disease or CAD. Coronary vasodilating capability is an independent prognostic determinant.

moderate to severe ischemia but with viable

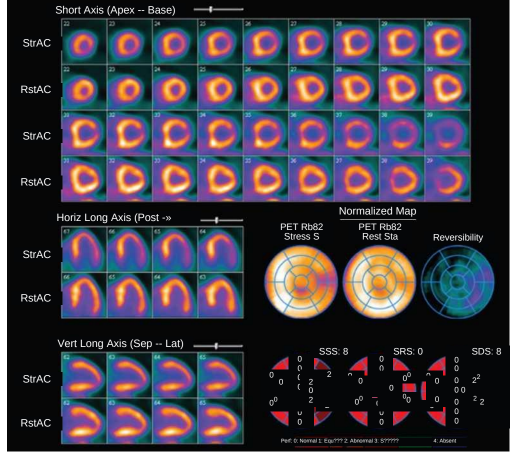


FIGURE 28-25 82Rb PET perfusion scan in a 62-year-old man with chest pain. The rest scans are normal, but there is a mild perfusion defect in the stress scan in the basal and mid anterolateral and inferolateral segments, consistent with reversible ischemia in the distribution of the left circumflex artery

TAE	TABLE 28—8 Stages of Cardiovascular Ischemia Detected by PET Scan					
	Rest Perfusion	Stress Perfusion	Contraction	Coronary Arteries	Metabolism	Diagnosis
1	Normal	Normal	Normal	Normal	Normal	Normal
2	Normal	Decreased	Normal	Normal	Normal	Endothelial or microvascular dysfunction Undetected atherosclerosis
3	Normal	Decreased	Normal	Stenosis	Normal	Ischemia by coronary artery stenosis
4	Decreased	Decreased	Decreased	Stenosis	Absent	Infarcted myocardium
5	Present	Decreased	Decreased	Stenosis	Present	Stunned myocardium
6	Decreased	Decreased	Decreased	Stenosis	Present	Hibernating myocardium

resulting in impaired contraction, which is persistent and results in perfusion-metabolism or prolonged transient ischemic episodes followed by reperfusion, suggests stunned myocar dium (Table 28-8, Fig. 28-26). resulting in depressed resting function with normal perfusion. Hibernating myocar dium is seen after repetitive ischemic episodes predictive values of 93%, 58%, 71%, and 86%, resulting in hypoperfusion at rest with depressed myocardial function. Both stunned and hibernating myocardium can recover function after revascularization.

In ischemia, the myocyte metabolism shifts from fatty acids to glucose. Therefore, uptake of an ¹⁸ F-labeled glucose analogue (FDG) in a region of myocardium indicates metabolic activity and

myocardium, the subset of patients who will benefit from thus viability. This enables PET to differentiate hibernating and revascularization. With prolonged imbalance between myocardial stunned myocardium from infarction. The presence of enhanced supply and demand, high-energy phosphates are depleted, FDG uptake in a region of decreased flow (known as a PET hibernating mismatch) indicates cell death or infarction. In stunned and hibernating myocardium, myocardium, whereas a reduction in both the metabolism and flow myocardial function is depressed at rest, but the myocytes are (perfusion-metabolism match) reflects nonviable infarcted viable. Stunned myocardium is seen after single or multiple, brief myocardium. Dysfunction with normal perfusion and metabolism

PET has mean sensitivity, specificity, and positive and negative

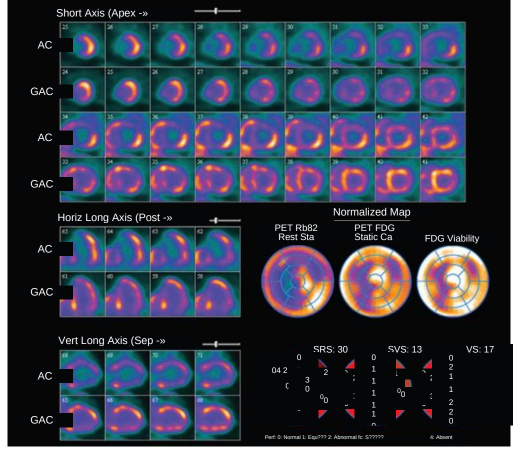


FIGURE 28-26 PET scan in advanced disease. In the rest perfusion images, reduced tracer uptake is seen in the basal and mid anterolateral, mid and apical anterior, inferior, and septal segments and the apex. In the FDG metabolic scan, there is unmatched FDG uptake in the basal anterolateral wall, mid and apical anterior and inferior segments, apex, mid inferior septum, and basal anterolateral wall, indicating hibernating segments in left anterior descending and right coronary artery territories (40% of myocardium). There is no FDG uptake in the mid anterior septum, anterolateral segments, and apical septum, indicating scar (18% of

respectively, to detect myocardial viability. 139 Infarcted in targeted imaging of atherosclerosis with molecular probes and is an independent predictor of improvement in ejection fraction novel drugs. after revascularization. 141 PET may be superior to SPECT in the setting of very severe left ventricular dysfunction. 142

Thus, PET is a useful noninvasive imaging modality to detect abnormal myocardial blood flow and MPR, which are reliable markers for disease progression and new endpoints of treatment aimed at improving global vascular function. Myocardial perfusion abnormalities in PET have been shown to improve after long-term intense risk factor modification.

CONCLUSION

MRI and PET are useful imaging techniques in the evaluation of different stages of atherosclerosis and assessment of cardiovascular risk factors. High-resolution multicontrast MRI is a valuable noninvasive technique in plaque characterization and quantification. Molecular MRI can detect subclinical stages of atherosclerosis. MRI is also useful in the evaluation of global and regional myocardial function, myocardial ischemia, and various vascular factors. Whole-body MRA can be used to evaluate multiple vascular beds in a single examination, particularly in the asymptomatic intermediate-risk group. PET scanning can be used

myocardium has only a 20% chance of functional improvement also in the evaluation of myocardial ischemia. Both MRI and PET after revascularization, compared with an 80% to 85% chance for are useful in the assessment of plaque progression or regression, hibernating or stunned myocardium. 140 The scar size on FDG-PET particularly as a surrogate endpoint of treatment in response to

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CHAPTER 29

Exercise Treadmill Stress Testing With and Without Imaging

Amil M. Shah and Samia Mora

KEY POINTS

- Exercise stress testing is a safe procedure in asymptomatic or minimally symptomatic individuals and provides both diagnostic and prognostic information.
- A growing body of literature supports the utility of nonelectrocardiographic parameters, including exercise capacity, chronotropic response, exercise blood pressure, and heart rate recovery, in determining prognosis in asymptomatic subjects beyond traditional risk factors or global risk scores.
- There are no Class I indications for exercise testing in asymptomatic individuals in the American College of Cardiology/American Heart Association guidelines because of insufficient clinical trial evidence of efficacy.
- Exercise testing may be considered in asymptomatic subjects before vigorous exercise is started if they are diabetic, older, at higher risk for CHD because of comorbid conditions, or involved in occupations potentially affecting public safety.
- Exercise testing with imaging should be used as an initial test in the following situations: preexcitation syndrome (Wolff-Parkinson-White), electrically paced ventricular rhythm, more than 1 mm of resting ST-segment depression, and complete left bundle branch block.
- Although some studies suggest additional prognostic value of imaging along with stress testing in asymptomatic individuals, this is not cost-effective in low-risk populations (low event rates and low specificity).

Data from 2006 indicate that 16,800,000 US adults aged 20 years or older have coronary heart disease (CHD), with an annual incidence of myocardial infarction (MI) of 1,450,000? Among patients experiencing sudden death, it is the first presentation of CHD in about 25%. ² For these reasons, there has been considerable interest in risk stratification of asymptomatic individuals without a diagnosis of CHD to prevent future events. ³ Given its noninvasive nature and the prognostic information it provides in established CHD, exercise testing has generated considerable interest.

A promising avenue for improving cardiac risk stratification has come from studies evaluating the prognostic value of exercise testing in asymptomatic populations with test variables that are not related to exercise-induced ST-segment depression. In particular, functional measures such as exercise capacity, blood pressure, and heart rate recovery have been linked to increased CHD and all-cause death in both women and men.

Additional imaging with myocardial perfusion or echocardiography may further identify another subgroup of asymptomatic individuals at higher risk for future events. However, despite the increased risk associated with certain aspects of the exercise test, the low cardiac event rate and low positive predictive value of abnormal test results in asymptomatic populations do not support a strategy of routine screening of adult asymptomatic populations. Clinical trial data are scarce and are eagerly awaited to answer the important question facing clinicians and patients alike: to screen or not to screen asymptomatic adults for CHD?

CURRENT GUIDELINES AND LIMITATIONS

Current guidelines do not recommend exercise testing for routine screening in asymptomatic subjects. Recent guidelines from the US Preventive Services Task Force found insufficient evidence to recommend routine screening because of the low positive predictive value (estimates ranging from 6% to 48%) among asymptomatic men. ⁴ Similarly, there are no Class I indications for exercise testing in asymptomatic adults in the 2002 American College of Cardiology/American Heart Association (ACC/AHA) guidelines (Table 29-1). However, these guidelines have been based largely on data regarding the performance of stress electrocardiography for diagnosis of coronary disease in a low-risk population. ⁵

There is a growing body of literature suggesting the utility nonelectrocardiographic exercise parameters in determining prognosis in asymptomatic subjects, beyond current risk stratification with commonly used global risk scores. 5 As reviewed in more detail later, these parameters include exercise capacity, chrono tropic response, exercise blood pressure, heart rate recovery, and ventricular arrhythmia . All have demonstrated prognostic utility even after accounting for traditional risk factors or global risk scores (eg, the Framingham risk

USE AS A SCREENING TEST

Broadly conceived, the purpose of a screening test is either earlier diagnosis of disease or risk stratification to allow effective interventions to prevent adverse outcomes. Current guidelines recommend office-based risk stratification of all individuals with multiple risk factor scores to determine global risk. 6 This is most commonly accomplished by the Framingham risk score as modified by the National Education Program Cholesterol Treatment Panel III. In this framework, patients can be stratified into low-risk (predicted 10-year absolute risk of MI or CHD death < 10%), intermediate risk (6% to 20%), and high-risk (> 20%) groups. 6

Whereas aggressive medical interventions are clearly indicated in high-risk patients, population-based studies suggest that less than 3% of asymptomatic subjects fall into the high-risk group, particularly

among women. ⁷ Although all patients should control known risk factors, some have argued that given the substantial variation in risk among intermediate-risk patients and the sizable proportion of patients falling in this category, these patients would benefit from further risk stratification (Fig. 29-1).

As presented in the following sections, multiple stress testing parameters have been demonstrated to add prognosis



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Class I None TABLE 29—1 2002 ACC/AHA Guidelines for the Use of Exercise Testing in Asymptomatic Individuals Without Known Coronary Heart Disease

Class IIa

Evaluation of asymptomatic persons with diabetes mellitus who plan to start vigorous exercise (Level of Evidence: C)

Class IIb

Evaluation of persons with multiple risk factors as a guide to risk reduction therapy

Evaluation of asymptomatic men older than 45 years and women older than 55 years:

Who plans to start vigorous exercise (especially if sedentary), or

Who are involved in occupations in which impairment might impact public safety, or

Who are at high risk for CAD due to other diseases (eg, peripheral vascular disease
and chronic renal failure)

Class III

Routine screening of asymptomatic men or women

From Gibbons RJ, Balady GJ, Bricker JT, et al: ACC/AHA 2002 guideline update for exercise testing: summary article: a report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines (Committee to Update the 1997 Exercise Testing Guidelines), Circulation 106:1883, 2002. © 2002 American Heart Association, Inc.

information beyond risk factor scores. However, data are still lacking that such refinements in prognostic assessment actually influence patient management and outcomes. ⁵ As discussed more fully in a later section, this critical information still awaits large-scale randomized clinical trials to assess the impact of risk stratification by exercise testing on outcomes. ⁵

EXERCISE STRESS TEST PERFORMANCE

Safety, Contraindications, and

Indications for Test Termination

Exercise stress testing is a generally safe procedure. Recognized serious complications of exercise testing include MI, malignant ventricular arrhythmias, and sudden death (Table 29-2). Large survey studies have reported acute MI in 0.9 to 3.6 per 10,000 tests, serious arrhythmias in 0.3 to 4.8 per 10,000 tests, and death in 0 to 0.5 per 10,000 tests. 8-10 The risk of adverse events is higher in post-MI patients and patients undergoing evaluation for malignant ventricular arrhythmias. ¹¹ Given the potential for serious risks (although rare), clinical judgment is essential in selecting patients appropriate for stress testing, as is careful monitoring by appropriately trained staff before, during, and after testing. ¹²

Absolute and relative contraindications to exercise testing are listed in Table 29-3. ¹³ In general, any patient with evidence of clinical or hemodynamic instability should not undergo exercise testing until the condition is stabilized. Absolute and relative indications for termination of exercise testing are listed in Table 29-4. ¹¹

Commonly Used Exercise Protocols

In general, exercise protocols are designed to assess exercise capacity. Maximum oxygen consumption ($V_{\circ,2}$ max), defined

Coronary heart disease risk assessment in asymptomatic patients:
Selective use of noninvasive testing following office-based risk assessment

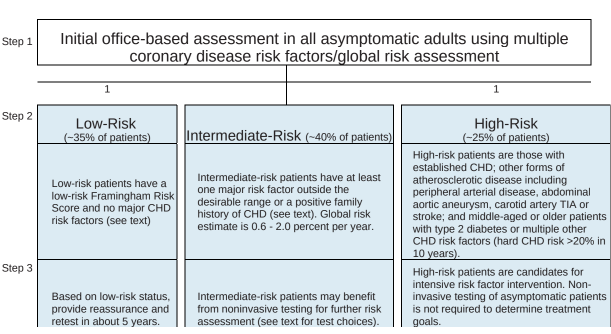


FIGURE 29-1 Potential role of exercise stress testing for further risk stratification of patients with intermediate risk of CHD based on global risk assessment. See text for further discussion. (From Greenland P Smith SC, Grundy SM: Improving coronary heart disease risk assessment in asymptomatic people: role of traditional risk factors and noninvasive cardiovascular tests. Circulation 104:1863, 2001.)

TABLE 29—2 Recognized Serious Complications of Exercise Testing

Cardiac

Bradyarrhythmias

Tachyarrhythmias

Acute coronary syndromes

Heart failure

Hypotension, syncope, and shock

Death

Noncardiac

Musculoskeletal trauma Soft tissue injury

Miscellaneous

Severe fatigue (malaise), sometimes persisting for days; dizziness; fainting; body aches; delayed feelings of illness

From Fletcher GF, Balady GJ, Amsterdam EA, et al: Exercise standards for testing and training: a statement for healthcare professionals from the American Heart Association. *Circulation* 104:1694, 2001. © 2001 American Heart Association, Inc.

TABLE 29—3 Absolute and Relative Contraindications to Exercise Stress Testing

absolute

Acute myocardial infarction (within 2 days)

High-risk unstable angina (as defined in the ACC/AHA Guidelines for the Management of Patients with Unstable Angina/Non-ST-Segment Elevation Myocardial Infarction)

Uncontrolled cardiac arrhythmias causing symptoms or hemodynamic compromise

Symptomatic severe aortic stenosis

Uncontrolled symptomatic heart failure

Acute pulmonary embolus or pulmonary infarction

Acute myocarditis or pericarditis

Acute aortic dissection

relationship

Left main coronary stenosis

Moderate stenotic valvular heart disease

Electrolyte abnormalities

Severe arterial hypertension (systolic blood pressure > 200 mm Hg and/or diastolic blood pressure > 110 mm Hg)

Tachyarrhythmias or bradyarrhythmias

Hypertrophic cardiomyopathy and other forms of outflow tract obstruction

Mental or physical impairment leading to inability to exercise adequately High-degree

atrioventricular block

From Gibbons RJ, Balady GJ, Bricker JT, et al: ACC/AHA 2002 guideline update for exercise testing: summary article: a report of the American College of Cardiology/ American Heart Association Task Force on Practice Guidelines (Committee to Update the 1997 Exercise Testing Guidelines). *Circulation* 106:1883, 2002. © 2002 American Heart Association, Inc.

as the maximum amount of oxygen a subject can take in from inspired air during dynamic exercise, is considered the best measure of cardiovascular fitness and exercise capacity. 11 When peak consumption is achieved, V O 2 max can estimate cardiac output. Oxygen uptake can be expressed in units of sitting/resting requirements or metabolic equivalents (METs); a MET is defined as a unit of sitting/resting oxygen uptake (approximately 3.5 mL O $_2$ /kg/min). "V $_{\rm 0.2}$ max varies significantly with age (declining by 8% to 10% per decade 14), gender (generally lower in women), physical activity (nearly 25% reduction noted with 3 weeks of bed rest 11), heredity, 15 and degree of myocardial impairment.

Exercise protocols with progressive incremental increases in workload tend to estimate "V O 2 max more accurately. ¹⁶ The optimal protocol will vary by patient and should last for 6 to

TABLE 29—4 Absolute and Relative Indications for Termination of an Exercise Stress Test

absolute

Drop in systolic blood pressure of > 10 mm Hg from baseline blood pressure despite an increase in workload, when accompanied by other evidence of ischemia

Moderate to severe angina

Increasing nervous system symptoms (eg, ataxia, dizziness, or near-syncope) Signs of poor perfusion (cyanosis or pallor)

Technical difficulties in monitoring ECG or systolic blood pressure Subject's desire to stop

Sustained ventricular tachycardia

ST elevation (> 1.0 mm) in leads without diagnostic Q waves (other than or aVR)

relationship

Drop in systolic blood pressure of > 10 mm Hg from baseline blood pressure despite an increase in workload, in the absence of other evidence of ischemia

ST or QRS changes such as excessive ST depression (> 2 mm of horizontal or downsloping ST-segment depression) or marked axis shift

Arrhythmias other than sustained ventricular tachycardia, including multifocal PVCs, triplets of PVCs, supraventricular tachycardia, heart block, or bradyarrhythmias

Fatigue, shortness of breath, wheezing, leg cramps, or claudication

Development of bundle branch block or IVCD that cannot be distinguished from ventricular tachycardia

Increasing chest pain

Hypertensive response (systolic blood pressure > 250 mm Hg and/or diastolic blood pressure > 115 mm Hg)

ECG, electrocardiogram; IVCD, intraventricular conduction delay; PVCs, premature ventricular contractions.

From Gibbons RJ, Balady GJ, Bricker JT, et al: ACC/AHA 2002 guideline update for exercise testing: summary article: a report of the American College of Cardiology/ American Heart Association Task Force on Practice Guidelines (Committee to Update the 1997 Exercise Testing Guidelines). Circulation 106:1883, 2002.
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12 minutes to reliably reflect the upper limit of the patient's cardiorespiratory function. 11,16 Exercise is most commonly performed with bicycle ergometry or treadmill. Commonly used protocols are illustrated in Figure 29-2. Compared with the cycle ergometer, treadmill tests tend to demonstrate 10% to 15% higher "V $_{\circ,}$ max , 5% to 20% higher peak heart rate, and more frequent ST-segment changes. 16

The most commonly used treadmill protocol in the United States is the Bruce protocol. ^{8,9} Whereas a large amount of published data exist with use of the Bruce protocol, the relatively large increments in work between stages can make "V O 2 max estimation less accurate and cause some patients to terminate exercise before "V O 2 max is achieved. ^{11,16} Estimation of V O 2 max appears more accurate with use of exercise duration-targeted ramp protocols, which constantly increase work by increasing incline at set brief intervals and increasing ramp speed on the basis of estimated functional capacity. ¹⁷ The major limitation is the need to accurately predict a patient's functional capacity.

STRESS TEST EXERCISES INTERPRETATION

Exercise testing produces both electrocardiographic and nonelectrocardiographic data that can be used both for diagnosis of CHD and for prognosis. Interpretation of exercise testing data must incorporate the clinical context of the test and, most importantly, the pretest probability of disease.



FIGURE 29-2 Commonly used treadmill and bike exercise protocols with the relationship between METs achieved and protocol stage. kpm, kilopond-meters; MPH, miles per hour; %GR, percent grade. (From Fletcher GF, Balady GJ, Amsterdam EA, et al: Exercise standards for testing and training: a statement for healthcare professionals from the American Heart Association. Circulation 104:1694, 2001. © 2001 American Heart Association, Inc.)

Performance Characteristics of Exercise Testing and Bayes' Theorem

Definitions of parameters used to quantify the diagnostic accuracy of a test are listed in Table 29-5. ¹³ Sensitivity defines the probability that a patient with disease will have a positive test result, and specificity defines the probability that a patient without disease will have a negative test result. However, the clinically relevant information in interpreting the results of any given test is the likelihood that a positive result is truly indicative of disease (positive predictive value) and that a negative result truly excludes disease (negative predictive value). As Table 29-5 illustrates, these parameters are dependent not only on the test but also on the prevalence of disease in the population (ie, the pretest probability).

Bayes' theorem states that the probability of disease after a diagnostic test is equal to the pretest probability of disease multiplied by the probability of a true positive result from the test. ¹³ A corollary is that the chances of a positive result truly reflecting disease (ie, positive predictive value) will be higher in high-prevalence populations and lower in low-prevalence populations. This point is illustrated in Figure 29-3. Bayesian analysis is critical in the appropriate interpretation of stress testing.

Test Interpretation: Diagnosis

Estimates of the diagnostic accuracy of electrocardiographic exercise stress testing for the diagnosis of hemodynamically significant coronary disease vary widely and are confounded

TABLE 29-5	
	Definitions of Sensitivity, Specificity, Positive Predictive Value, and Predictive Accuracy
Sensitivity	[TP/(TP + FN)] x 100
Specificity	[TN/(FP + TN)] x 100
Positive predicted value	Sensitivity X P(CAD) [Sensitivity X P(CAD)] + [(1 — specificity)[1 — P (CAD)]]
Predictive accuracy	[Sensitivity x P(CAD)] + [(1 — specificity)[1 - P(CAD)]]

Note that the calculated positive predictive value (PPV) and negative predictive value (NPV) are dependent on the population prevalence of disease. FN, false negatives; FP, false positives; P(CAD), pretest probability; TN, true negative; TP, true positive. From Gibbons RJ, Balady GJ, Bricker JT, et al: ACC/AHA 2002 guideline update for exercise testing: summary article: a report of the American College of Cardiology/ American Heart Association Task Force on Practice Guidelines (Committee to Update the 1997 Exercise Testing Guidelines). Circulation 106:1883, 2002. © 2002 American Heart Association, Inc.

by the fact that the majority of studies suffer from workup bias, discussed in further detail later.

Determining the Pretest Probability of CHD

Multiple predictive models have been developed to assist the clinician in assessing the pretest probability of CHD in a given patient. ^{13,18-20} These models consistently show age, gender, and chest pain history to be the most powerful

Interpretation of Electrocardiographic Response

Normally encountered electrocardiographic changes with exercise include increased P wave magnitude in the inferior leads with shortening of the PR interval, decreased R wave amplitude in the lateral leads, and depression of the I-point in the lateral leads. 11 Assessment of electrocardiographic manifestations of exercise-induced myocardial ischemia focuses on the ST segment.

In exercise electrocardiography, the ST-segment deviation is measured relative to the PQ junction. 11 Generally accepted criteria for abnormal ST-segment depression are horizontal and downsloping ST-segment depression of > 0.10 mV (1 mm) for 80 msec, with downsloping ST-segment depression being more specific than horizontal or upsloping ST-segment depression (Fig. 29-4). 11

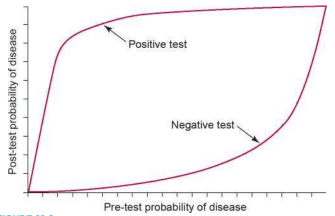


FIGURE 29-3 Graphic illustration of the post-test probability of disease (y -axis) for a positive versus negative test as a function of the pretest probability of disease (X -axis). (Modified from Schwartz JS: Clinical decision-making in cardiology. In Zipes DP, Libby P Bonow RO, Braunwald E, editors: Braunwald's heart disease: a textbook of cardiovascular medicine, ed 7, Philadelphia, Elsevier Saunders, 2005, pp 27-34.)

ST-Segment Depression. Resting ST-segment depression is a risk marker of adverse cardiac prognosis in itself. 21 Resting STsegment depression of < 1 mm has been shown to increase the sensitivity but to decrease the specificity of exercise testing, ^{22,23} and exercise testing is still considered a reasonable first test in these patients. ¹³ Importantly, ischemic ST-segment depression occurring only during the recovery phase of an exercise test appears to have comparable diagnostic significance to STsegment depression occurring during exercise. 24

Exercise-induced ST-segment depression has diagnostic properties that vary widely. Important methodological limitations that may inflate estimates of ST-segment depression sensitivity are the inclusion of subjects with high probability of having disease (eg, prior MI) and workup bias. ¹³ Workup bias refers to the inclusion of subjects based on the results of the test 29 being evaluated, that is, only subjects undergoing both stress testing and coronary angiography are included, although the decision to pursue angiography is influenced by the results of the exercise test. 13

A meta-analysis of 147 studies involving 24,074 patients and comparing exercise-induced ST depression with coronary angiography reported a mean sensitivity and specificity of 68% and 77%, respectively. 25 However, both the sensitivity and specificity calculations varied widely; the range of reported sensitivity was 23% to 100%, and the range of reported specificity was 17% to 100%. 25 Studies that avoided workup bias and did not include many patients with high pretest probability of disease suggest a sensitivity of 50% and a specificity of 90% associated with exercise-induced 1-mm ST-segment depression.

ST-Segment Elevation. The development of ST-segment elevation, measured from the baseline ST level, is not infrequent in leads with preexisting Q waves and is of unclear significance among patients with prior MI. 13 Exercise-induced ST-segment elevation in subjects without preexisting Q waves is rare, occurring in an estimated 0.1% of patients in a clinical laboratory. It is associated with transmural ischemia and reliably localizes the area of ischemia. 13

Gender Differences in ST-Segment Changes. It is notable that exercise-induced ST-segment depression has not been found to be associated with higher risk in asymptomatic women, ²⁶⁻²⁸ in contrast to findings in asymptomatic men, in whom ischemic electrocardiographic changes have been associated with higher mortality. 29-32 This sex difference in the prognostic accuracy of the ST segment is consistent with previously reported sex differences regarding its diagnostic accuracy 33 and may be related to hormonal effects on the electrocardiogram 34 or sex differences in endothelial function. 35 Other measures obtained from exercise testing that add

TABLE 29-6		ly Used Table Incorporating Age, Gender,	and Character of Chest Pain to Estimate Pre	test Probability of Coronary He	art Disease
Age (years)	Gender	Typical/Definite Angina Pectoris	Atypical/Probable Angina Pectoris	Nonanginal Chest Pain	Asymptomatic
30-39	Men	Intermediate	Intermediate	Low	Very low
	Women	Intermediate	Very low	Very low	Very low
40-49	Men	^{Hi} g ^h	Intermediate	Intermediate	Low
	Women	Intermediate	Low	Very low	Very low
50-59	Men	^{Hi} g ^h	Intermediate	Intermediate	Low
	Women	Intermediate	Intermediate	Low	Very low
60-69	Men	Hi g h	Intermediate	Intermediate	Low
	Women	Hi g h	Intermediate	Intermediate	Low

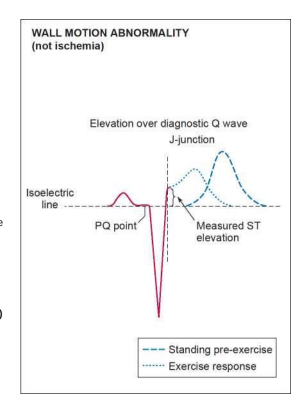
See text for further discussion

From Gibbons RJ, Balady GJ, Bricker JT, et al: ACC/AHA 2002 guideline update for exercise testing: summary article: a report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines (Committee to Update the 1997 Exercise Testing Guidelines). Circulation 106:1883, 2002. © 2002 American Heart Association, Inc

Flat or downsloping deviation greater than or equal to 1.0 mm is abnormal

Slope is determined over 60 msec (1.5 boxes)

ST deviation amplitude is relative to the isoelectric line and measured at the Jiunction Flat Downsloping (worse)



BORDERLINE

Flat or downward sloping

Upsloping

Flat or downsloping ST deviation greater than or equal to 0.5 mm and less than 1.0 mm is borderline

Slope is determined over 60 msec (1.5 boxes)

ST deviation amplitude is relative to the isoelectric line and measured at the J-junction

Flat

Downsloping (worse)

Isoelectric line Upsloping depression with ST60 greater than or equal to 2.0 mm is borderline

(1.5 boxes)

Notes: To be borderline, ST depression at ST0 must be greater than 2 mm also

FIGURE 29-4 Standard interpretation of ST-segment

deviation for exercise electrocardiography. (From Fletcher GF, Balady GJ, Amsterdam EA, et al: Exercise standards for testing and training: a statement for healthcare professionals from the American Heart Association. Circulation 104:1694, 2001. © 2001 American Heart Association, Inc.)

useful prognostic information in women is discussed later and includes functional capacity and heart rate recovery.

Test Interpretation: Prognosis

Use of Common Prognostic Scores (Duke Treadmill Score)

Multiple parameters measured during the exercise stress test yield prognostic information. Many are discussed in greater detail later. Multiple studies have demonstrated the prognostic importance of exercise capacity, measured as treadmill stage, exercise duration, metabolic equivalents, watts, or double product. 13 In addition, studies evaluating the most predictive independent prognostic parameters from exercise tests

consistently also identify the presence of exercise-induced myocardial ischemia, generally reflected in ST-segment deviation or anginal symptoms. 13 Whereas

Progn	Duration of exercise		
5-yea survivalr		MET 20	Min 18
i.	0.2% 0.4%	17	15
		13	12
0.93	1% 1.5%	10	9
	2% 3%	7	6
0.80	1% 5%	5	3
070- 0.55	6% 9%	0	0



2 mm

ST-segment deviation

during

mm-i-

1 mm

exercise 0

Ischemia-

reading

line

3mm

multiple risk scores integrating these various prognostic markers have been developed, ¹³ the most widely used is the Duke

Exercise capacity is the most powerful prognostic parameter from an exercise test. Multiple large studies in both asymptomatic and symptomatic subjects have reported similar findings. ^{39,40-44} This relationship is present in both men ^{31,39} and

4 mm-1-

FIGURE 29-5 Nomogram to predict prognosis (annual mortality rate and 5-year survival) based on the Duke treadmill score. (From Gibbons RJ, Balady GJ, Bricker JT, et al: ACC/AHA 2002 guideline update for exercise testing: summary article: a report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines [Committee to Update the 1997 Exercise Testing Guidelines]. Circulation 106:1883, 2002. © 2002 American Heart Association, Inc.)

Angina

during

exercise -rNone

Nonlimiting

Exerciselimiting

treadmill score (DTS). 36,37

The DTS was initially developed in 2842 subjects referred for cardiac catheterization who had also undergone exercise testing for evaluation of symptoms of CHD. 36 The investigators identified three prognostic variables: ST-segment depression , exercise time on Bruce protocol, and Duke angina index. In the Duke angina index, angina index 0 = no angina, 1 = typical angina occurred, and 2 = angina was reason for test termination.

The resulting DTS is calculated as

DTS = Exercise time (in minutes) - (5 x ST deviation in mm)

- (4 x Duke angina index)

This score effectively stratified subjects in terms of 5-year risk of death or MI and provided incremental prognostic value beyond clinical and angiographic data. In subjects with low-risk treadmill scores (>+ 5), 5-year event-free survival (death or MI) was 93%; in intermediate-risk subjects (score - 10 to + 4), it was 86%; and in high-risk subjects (score < - 11), it was 63%. 36 The investigators subsequently validated the score in unselected outpatients, demonstrating that in this population , the 4-year survival among low-risk patients was 99%, whereas among high-risk patients, it was 79% (Fig. 29-5). 37 An important limitation of the DTS is its limited discrimination in elderly subjects. 38

Nonelectrocardiographic Prognostic Parameters

Exercise Capacity. Exercise (or functional) capacity refers to the maximal oxygen extraction obtainable during exercise and is commonly measured in METs. METs have multiples of basal metabolism; one MET is the basal oxygen uptake during quiet sitting and is equal to 3.5 mL/kg/min. Exercise capacity is influenced by factors outside of cardiovascular fitness, most importantly age and gender. ^{39,40} Nomograms have been developed to estimate age-predicted exercise capacity among men [18.0 - (0.15 x age)] ³⁹ and women [14.7 - (0.13 x age)] ⁴⁰ (Fig. 29-6). For example, with the use of this nomogram, women who did not achieve 85% of their age-predicted exercise capacity had a twofold higher risk of cardiovascular death. ⁴⁰

women. 26,40 In one landmark study of more than 14,000 healthy men and women, physical fitness measured by maximal treadmill time was significantly associated with mortality during 8-year follow-up, independent of demo graphics and standard risk factors. 45

A study of 2994 asymptomatic women observed for 20 years demonstrated that women who were below the median at baseline for exercise capacity (< 7.5 METs) and heart rate recovery (< 55 beats/min difference between peak exercise heart rate and heart rate at 2-minute recovery) had a 3.5-fold higher risk of cardiovascular death independent of traditional risk factors. ²⁶ Importantly, functional capacity provides prognostic information beyond traditional risk stratification by the Framingham risk score in both men ²⁷ and women, ⁴⁶ as discussed later.

Chronotropic Incompetence. Chronotropic incompetence refers to an inability to achieve the expected increase in heart rate with exercise. ⁵ Multiple parameters have been used to assess chronotropic incompetence. It is most commonly evaluated by the proportion of age-predicted maximal heart rate (HR) achieved during the stress test (peak HR/220 - age). ⁵ However, in addition to age, the chronotropic response to exercise is also affected by resting heart rate and physical fitness. ⁴⁷ The proportion of heart rate reserve used is defined as [(peak HR rest HR)/(maximum age-predicted HR - rest HR)] x 100 and incorporates information about resting heart rate. ⁴⁸ The chronotropic index incorporates data for both

PERCENTAGE OF PREDICTED EXERCISE CAPACITY FOR AGE

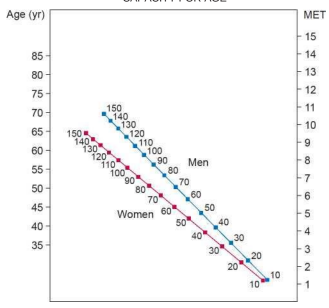


FIGURE 29-6 Nomograms for determination of age-predicted exercise capacity among men and women. (From Gulati M, Black HR, Shaw LJ, et al: The prognostic value of a nomogram for exercise capacity in women. N Engl J Med 353:468, 2005. © 2005 Massachusetts Medical Society. All rights reserved.)

resting heart rate and physical fitness and is defined as the ratio of the metabolic reserve to the heart rate reserve: 47

Chronotropic index at any stage of the exercise test =

[(MET S stage - METS rest) /(METS peak - METS

(HR peak - HR rest) $^{/\!(HR)}$ max predicted HR rest)

Chronotropic incompetence measured by all three indices has been consistently associated with increased risk of all causes and cardiovascular mortality, even after adjustment demographics and standard risk factors. 47-50 This has been shown in both referral populations 48,49,50 and studies of healthy asymptomatic individuals. 47 In studies involving nuclear perfusion imaging, measures of chronotropic incompetence consistently provide additional prognostic information beyond the findings on perfusion imaging. 48,49 In asymptomatic individuals, measures of chronotropic incompetence appear to provide additional prognostic information beyond the Framingham risk score. 27 Importantly, the majority of studies assessing the prognostic role of chronotropic incompetence excluded patients receiving beta blocker therapy at the time of the exercise test.

Mechanistically, the increase in heart rate with exercise is thought to reflect physiological parasympathetic withdrawal and increased sympathetic activity with exercise. ⁵ An early study comparing chronotropic response among subjects with heart failure and normal controls suggested that in this population at least, chronotropic incompetence is at least partially mediated by postsynaptic beta-adrenergic desensitization with resulting decreased sensitivity of the sinus node to sympathetic activity. ^{5,51}

Heart Rate Recovery. Whereas chronotropic response to exercise is thought to reflect sympathetic sensitivity, the rate of decrease in heart rate after exercise, termed heart rate recovery, likely reflects parasympathetic reactivation. ⁵

heart rate Heart rate recovery = heart rate heart rate peak **exercise**

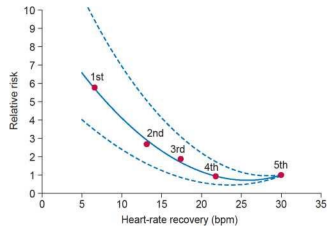


FIGURE 29-7 Relationship between heart rate recovery (beat per minute reduction during the first minute after exercise) and all-cause mortality at 6-year follow-up among a referral population of more than 2400 individuals. Relative risks for each quintile of heart rate recovery compared with the highest quintile (5th) are presented. (From Cole CR, Blackstone EH, Pashkow FJ, et al: Heart-rate recovery immediately after exercise as a predictor of mortality. N Engl J Med 341:1351, 1999. ©1999 Massachusetts Medical Society. All rights reserved.)

1 or 2 minutes after exercise

Heart rate recovery is a measure of parasympathetic nervous system function and autonomic balance. 52 Impaired heart rate recovery is commonly defined as a decrease in heart rate of < 12 beats/min within the first minute after exercise , $^{53-55}$ although some have proposed a decrease of < 22 beats/min at 2 minutes after exercise as the optimal cut point for prediction of risk of mortality. 56

Impaired heart rate recovery is associated with an increased risk of death, even after adjustment for patient demographics, standard risk factors, and perfusion abnormalities on nuclear imaging (Fig. 29-7). ⁵³⁻⁵⁵ The relationship between heart rate recovery and risk of death is also independent of exercise capacity and peak chronotropic response. ^{53,55} Importantly, impaired heart rate recovery, in combination with exercise capacity, provides incremental prognostic information beyond established global risk scores, including the Framing ham risk score ⁴⁶ and the European SCORE, ⁵⁷ as discussed in the next section.

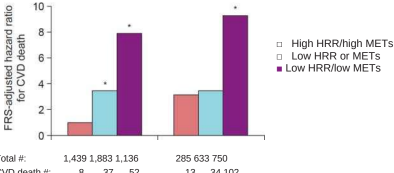
How concurrent use of beta blockers affects the predictive power of heart rate recovery is unclear. In multiple studies, subjects with abnormal heart rate recovery do appear more likely to be receiving beta blocker therapy. ^{54,55} However, sub group analysis in more than 2400 subjects referred for exercise testing, 13% of whom were receiving beta blocker therapy, did not find any significant modification of the relationship between impaired heart rate recovery and risk of death by concurrent beta blocker use. ⁵³

Incremental Value of Exercise Capacity and Heart Rate Recovery Beyond Traditional Risk Scores. Recently data - demonstrated that these two variables added important prognostic information to the Framingham risk score. Asymptomatic individuals with low- or intermediate-risk Framingham scores had 8- to 10-fold higher risk of cardiovascular death if they also had low values of these two exercise measures combined (Fig. 29-8). 46 Using annual cardiovascular mortality rates and receiver operating curves, the study found that half of women and just under half of men with Framingham risk scores of 10% to 19%, in addition to half of women with Framingham risk scores of 6% to 9%, would be reclassified as high risk on the basis of having low heart rate recovery/low METs, thus providing clinically important risk information.

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By applying the simple measurements of heart rate recovery and exercise capacity, this study accurately reclassified

FIGURE 29-8 Framingham risk score (FRS)-adjusted hazard ratios for cardiovascular death from Cox proportional hazards models according to FRS categories and heart rate recovery/metabolic equivalents (HRR/METs) groups at 20-year follow-up. F_{trend} values are for tests of significance for trend across the three HRR/METs groups in each FRS category. An asterisk is shown for each of the hazard ratios that were statistically significant in a pairwise comparison to individuals with high HRR/high METs in the same FRS group (all < 0.001). (From Mora S, Redberg RF, Sharrett AR, Blumenthal RS: Enhanced risk assessment in asymptomatic individuals with exercise testing and Framingham risk scores. Circulation 112:1566, 2005.)



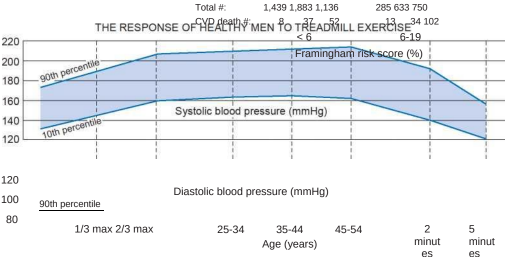


FIGURE 29-9 Illustration of the normal blood pressure response to exercise. (From Fletcher GF, Balady GJ, Amsterdam EA, et al: Exercise standards for testing and training: a statement for healthcare professionals from the American Heart Association. Circulation 104:1694, 2001. © 2001 American Heart Association, Inc.)

as high risk approximately 50% of asymptomatic individuals who pressure. 64,65 However, these findings were controversial because were deemed to be at intermediate risk by Framingham criteria. the relative contribution of rest blood pressure may have been These findings have been substantiated in other populations. 27,57,58 underestimated in these studies, and the findings were attenuated Finally, both exercise capacity and heart rate recovery are at least once subjects were asked to exercise to maximum capacity. Given partially modifiable and may be improved with moderate regular these conflicting data on the association of systolic blood pressure physical activity and exercise training by approximately 15% to 30% during exercise and cardiovascular endpoints, the ACC/AHA in a period of several months. 59

to exercise (Fig. 29-9) is characterized by a steady rise in systolic did not link it with cardiovascular disease or mortality outcomes. 13 blood pressure with little change in diastolic blood pressure, thereby resulting in an increase in pulse pressure (systolic minus cardiovascular endpoints remained unclear until recently. In 2008, diastolic blood pressure). 11 Exercise-induced hypotension is often investigators for the Framingham Offspring Study noted an defined as an initial increase in blood pressure followed by a association between low-level exercise diastolic blood pressure and decrease of 20 mm Hg during exercise or by a decrease in blood cardiovascular disease events in asymptomatic individuals pressure during exercise > 10 mm Hg below standing rest blood independent of traditional risk factors and rest blood pressure. 67 pressure. 11.60 It is associated with an increased prevalence of three- Recently, a study was performed that found a significant positive vessel disease or left main coronary artery disease and up to association for blood pressure threefold increased risk of death at 2-year follow-up. 60

Studies including normotensive men, women, whites, and African Americans have suggested that exaggerated systolic blood pressure response to exercise (commonly defined as peak systolic blood pressure > 200 to 220 mm Hg) is associated with an increased risk of subsequent hypertension. 61,62 However, studies have been conflicting.

Fagard and colleagues 63 first reported on this association in their study of 143 hypertensive men, demonstrating that blood pressure attained at submaximal or maximal workloads was not predictive of death from cardiovascular disease once baseline blood pressure was taken into account. Contrary to this, several studies showed that systolic exercise blood pressure could independently predict death from cardiovascular disease. 64-66 In particular, two landmark studies examining more than 7000 individuals found that submaximal exercise systolic blood pressure was, in fact, predictive of future cardiovascular death, even after controlling for rest blood

guidelines on exercise testing included exercise-induced Blood Pressure Responses. The normal blood pressure response hypertension as a marker for future clinical hypertension, but they

The relative contribution of diastolic blood pressure to

498 (both systolic and diastolic) at rest, low-level (Bruce stage 2) exercise, and maximum exercise with death from cardiovascular disease in an asymptomatic population consisting of more than 6500 individuals. 68 The strongest stage 2, and then maximal blood pressure during exercise. Among hypertensives, whether they had normal (< 120/80) or prehypertension, Bruce stage 2 blood pressure > 180/90 versus < 180/90 mm Hg carried a 1.5- to 2.0-fold increased risk of cardiovascular death, independent of rest blood pressure risk factors.

In addition, exercise blood pressure added predictive deserve further investigation.

Ventricular Arrhythmias. The prognostic role of exerciseinduced ventricular ectopy remains controversial, 11 with conflicting data. 69-71 Studies among population-based cohorts 72,73 and referral populations 74 suggest that exercise-induced frequent ventricular ectopy is associated with increased longterm all-cause and cardiac mortality. In these analyses, frequent ventricular ectopy is generally defined as increased frequency of premature ventricular contractions (eg, > 7 per tachycardia.

In a study of 6101 asymptomatic men who underwent exercise testing, exercise-induced frequent ventricular ectopy was associated with increased risk of cardiovascular death at 23-year follow-up. 72 This association was independent of and similar in magnitude to the relationship between exerciseinduced ST-segment depression and risk of cardiovascular death (2.5-fold increased risk). 72 Similarly, an analysis of 2885 asymptomatic subjects from the Framingham Offspring Study found that subjects with exercise-induced premature ventricular contractions were at increased risk of all-cause mortality at 15-year follow-up, with a multivariable adjusted hazard ratio of nearly twofold. 73 A recent report from a refer STRESS TESTING WITH IMAGING FOR ral population of 29,244 patients found that the presence of frequent ventricular ectopy during recovery phase was an even stronger predictor of mortality than frequent ectopy

ASYMPTOMATIC INDIVIDUALS during exercise at 5 years of follow-up. 74

but more data are needed to clarify this association. The strength and magnitude of this relationship likely vary with the definition of frequent ventricular ectopy employed, the phase of exercise test when measured (exercise versus recovery), the population being assessed (asymptomatic screening versus referral versus known CHD), and the duration of follow-up.

Silent Ischemia

Silent ischemia is defined as the presence of demonstrable myocardial ischemia in the absence of angina symptoms. have been done in subjects with established coronary disease, either prior treated angina or prior acute coronary syndrome. 75-77 In the Asymptomatic Cardiac Ischemia Pilot (ACIP) ischemia exercise testing and randomized to medical therapy (angina guided or ischemia screening. guided) or revascularization. 78

Among the medically treated subjects, the burden of ischemia reflected in the number of ischemic episodes on electrocardiographic monitoring significantly associated with the incidence of ischemic events at 1-year follow-up. 78 The presence of myocardial ischemia was also associated with increased risk of cardiac events after revascularization 75 and MI. 77

Fewer data exist for the prevalence and prognostic significance association existed for rest blood pressure, followed by Bruce of silent ischemia in asymptomatic subjects without known coronary disease. Ischemic electrocardiographic response to stress has been associated with increased risk of subsequent cardiac events but has poor predictive value because of the high false-positive rate in low-risk populations. 79 Studies of asymptomatic ischemia on exercise testing confirmed by angiography suggest a prevalence among otherwise healthy men of roughly 2.5% to 2.7%. 80,81

In a report of 407 subjects from the Baltimore Longitudinal Study value (net reclassification improvement, 10%-12%). These of Aging, investigators noted an age-dependent increase in the findings could potentially identify individuals who warrant 29 prevalence of concordant abnormal electrocardio graphic and more aggressive treatment than is currently recommended and nuclear perfusion responses to exercise stress testing, from 2% for subjects younger than 60 years to 15% for subjects older than 80

Even greater uncertainty exists about the prognostic implications of silent ischemia in asymptomatic populations. In the analysis from the Baltimore Longitudinal Study of Aging, during an average follow-up of 4.6 years, subjects with concordant electrocardiographic and perfusion abnormalities with stress testing were at a higher risk (48%) of subsequent angina, MI, or cardiac death than were those with normal results (7%) or with abnormal minute or > 10% of beats in a 30-second period), ventricular response of only one parameter (8%). ⁷⁹ Importantly, the majority of bigeminy or trigeminy, couplets or triplets, or ventricular events among those with concordant abnormalities consisted of the development of angina (6 of 11), and the number of subjects with concordant abnormalities was small (n = 23).

Results of studies using ambulatory electrocardiographic recording to detect asymptomatic ischemia have been conflicting. 82,83 In a study of 394 men born in 1914 from Sweden, 79 of 341 subjects without a history of CHD had at least one episode of horizontal or downsloping ST depression of at least 0.1 mV. During an average follow-up of 43 months, these 79 subjects were at a higher risk of fatal or nonfatal MI (10.1% versus 2.3%). 83 However, not all studies have found this association with hard outcomes.

SCREENING RISK ASSESSMENT OF

Current guidelines do not recommend routine screening with an Data exist that the occurrence of exercise-induced frequent exercise test for detection of CHD but do allow exercise testing in ventricular ectopy is associated with increased risk of death, individuals with risk factors before starting a vigorous exercise program other than walking (see Table 29-1). Whereas both exercise myocardial perfusion scintigraphy and exercise echocardiography have been shown to have prognostic value in symptomatic patients, their role for screening or prognostic purposes in asymptomatic individuals has not been well examined. Here, we summarize accepted indications for testing with imaging and examine the few prospective studies that have evaluated the use of stress imaging in asymptomatic or mildly symptomatic populations.

Although some studies, but not all, found additional prognostic value to stress-induced imaging abnormalities, absolute event rates Most investigations of the clinical relevance of silent ischemia were consistently low (even in subjects with abnormal test results), and the sensitivity was too low to justify cost-effective use of these tests. One randomized clinical trial, the Detection of Ischemia in Asymptomatic Diabetics (DIAD) study, was performed with study, 496 subjects with stable coronary disease, evidence of pharmacologic stress but is discussed in detail because it is the only ambulatory randomized clinical trial to test a strategy of screening stress electrocardiographic monitoring, and coronary disease myocardial perfusion imaging (MPI) versus no screening in amenable to revascularization by angiography were asymptomatic individuals and found no benefit in outcomes for

Indications for Testing with Imaging

In the diagnosis of CHD, exercise testing with imaging as an initial test is recommended only in a limited number of circumstances (Table 29-7). 84 These are situations in which base line electrocardiographic abnormalities make interpretation of ischemic electrocardiogram and abnormal thallium scan had a 3.6-fold ST-segment deviation unreliable: preexcitation syndrome (Wolff- increased risk of clinical CHD during a 5-year period. However, the Parkinson-White), electrically paced ven tricular rhythm, more than prevalence of ischemia was low as expected for a general 1 mm of resting ST-segment depression, and complete left bundle community-based population, and the authors concluded that it branch block. 13 Subjects with ventricular paced rhythms and complete left bundle branch block should generally undergo vasodilator stress perfusion studies because of the increased falseimaging. Subjects unable to exercise should undergo a pharmacological stress imaging study.

TABLE 29-7

2002 ACC/AHA Guidelines for the Use of

Imaging as a First-Line Test for the Diagnosis of Coronary Heart Disease in Patients Who Are Able to Exercise

Class I

Exercise myocardial perfusion imaging or exercise echocardiography in patients with an intermediate pretest probability of CHD who have one of the following baseline ECG abnormalities:

Preexcitation syndrome (Wolff-Parkinson-White) (Level of Evidence: B) More than 1-mm ST-segment depression at rest (Level of Evidence: B) Exercise

myocardial perfusion imaging or exercise echocardiography in patients with prior revascularization (PCI or CABG) (Level of Evidence: B)

Vasodilator (adenosine or dipyridamole) myocardial perfusion imaging in patients with an intermediate pretest probability of CAD and one of the following baseline ECG abnormalities:

Paced ventricular rhythm (Level of Evidence: C) Left bundle branch block (Level of Evidence: B)

Class IIb

Exercise myocardial perfusion imaging or exercise echocardiography in patients with a low or high probability of CHD who have one of the following baseline ECG abnormalities:

Preexcitation syndrome (Wolff-Parkinson-White) (Level of Evidence: B)

More than 1-mm ST-segment depression at rest (Level of Evidence: B) Vasodilator (adenosine or dipyridamole) myocardial perfusion imaging in

patients with a low or high pretest probability of CAD and one of the following baseline ECG abnormalities:

Paced ventricular rhythm (Level of Evidence: C)

Left bundle branch block (Level of Evidence: B)

Exercise myocardial perfusion imaging or exercise echocardiography in patients with an intermediate probability of CHD who have one of the following:

Digoxin use with less than 1-mm ST depression on the baseline ECG (Level of

LVH with less than 1-mm ST depression on the baseline ECG (Level of Evidence: B) Exercise myocardial perfusion imaging, exercise echocardiography, adenosine or dipyridamole myocardial perfusion imaging, or dobutamine echocardiography as the initial stress test in a patient with a normal rest ECG who is not taking digoxin (Level of Evidence: B) Exercise or dobutamine echocardiography in patients with left bundle branch block (Level of Evidence: C)

CABG, coronary artery bypass grafting; CAD, coronary artery disease; CHD, coronary heart disease; ECG, electrocardiogram; LVH, left ventricular hypertrophy; PCI, percutaneous

From Gibbons RJ, Balady GJ, Bricker JT, et al: ACC/AHA 2002 guideline update for exercise testing: summary article: a report of the American College of Cardiology/ American Heart Association Task Force on Practice Guidelines (Committee to Update the 1997 Exercise Testing Guidelines). Circulation 106:1883, 2002. © 2002 American Heart Association, Inc.

Myocardial Perfusion Imaging

The Baltimore Longitudinal Study of Aging evaluated the predictive value of exercise testing with imaging in a communitybased study. ⁷⁹ Individuals with an abnormal exercise was not cost-effective to screen unselected individuals, especially those younger than 60 years.

For higher risk asymptomatic patients enriched with risk factors positive rate associated with exercise stress and echocardiographic and having a family history of premature CHD, thallium imaging imaging. 84 Resting ST-segment depression < 1 mm does not require combined with exercise testing improved the predictive value of the test. 85 In the Johns Hopkins Sibling Study, 264 asymptomatic siblings (mean age, 46 years) of individuals with premature CHD 29 underwent exercise thallium testing. The relative risk for development of clinical CHD (death, MI, or revascularization) during a follow-up period of 6 years was fourfold for an abnormal exercise electrocardiogram and fivefold for an abnormal scan, defined as a moderate reversible segmental perfusion defect. Siblings with an abnormal exercise electrocardiogram and abnormal thallium scan had a relative risk of 14.5 compared with siblings who had both normal. In a subsequent report from the same study but with a larger sample size of 734 siblings, abnormal thallium scan combined with abnormal exercise electrocardiogram correlated with generally mild obstructive coronary disease on angiography. 86

The DIAD Study

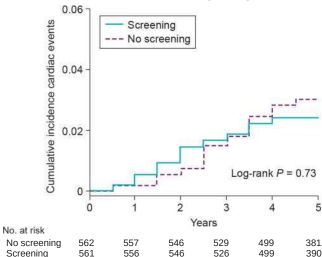
The primary objective of the DIAD study was to determine whether screening of type 2 asymptomatic diabetic patients with pharmacological (adenosine) MPI would reduce cardiac events, defined as fatal and nonfatal coronary events. 87 A total of 1123 type 2 diabetic patients with no history of CHD (mean age, 60 years; mean diabetes duration, 8 to 9 years; > 60% with two or more additional cardiac risk factors at baseline) were randomized to screening with adenosine MPI versus no screening. Patients were then referred to their modified then referred to their medical providers for appropriate care without mandating angiography in those with abnormal stress test results.

During a mean follow-up of 5 years, the primary endpoint was not significantly different (P = 0.73; Fig. 29-10A) among those screened with MPI (2.7%) versus those not screened (3.0%), despite a higher use of angiography in the MPI compared with the nonscreened group (4.4% versus 0.5%). Overall, the cardiac event rate during follow-up was low (0.6%/year), with most patients receiving statins and aspirin. This low cardiac event rate in this asymptomatic diabetic population is lower by up to fourfold compared with diabetic patients referred for stress testing (workup/referral bias).

Whereas moderate or large perfusion abnormalities on MPI had sixfold higher relative risk of cardiac events compared with normal studies or small perfusion abnormalities (Fig. 29-10B), the positive predictive value was low (12%), with more than half of the cardiac events occurring in patients with normal studies. Thus, the DIAD study found no clinical benefit during a 5-year period for routine screening of type 2 diabetic patients without history of CAD with adenosine MPI, despite the higher risk associated with moderate to large perfusion abnormalities, calling into question certain guide line recommendations for screening of asymptomatic individuals (see Table 29-1). 13

Exercise Echocardiography

The predictive value of exercise echocardiography was evaluated in 1859 individuals (mean age, 51 years) with no angina



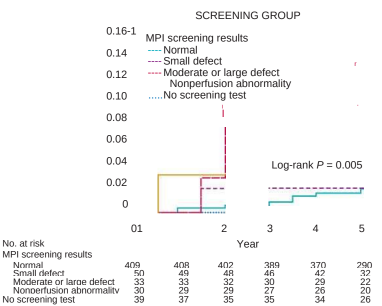


FIGURE 29-10 Cumulative incidence of cardiac events in asymptomatic type 2 diabetics enrolled in the DIAD trial. A. Subjects randomized to screening stress myocardial perfusion imaging (n = 561) compared with no screening (n = 562). B, Cumulative incidence among subjects randomized to screening by myocardial perfusion imaging results. (From Young LH, Wackers FJT, Chyun DA, et al: Cardiac outcomes after screening for asymptomatic coronary artery disease in patients with type 2 diabetes: the DIAD study: a randomized controlled trial. JAMA 301:1547, 2009. © 2009 American Medical Association. All rights reserved.)

or heart failure symptoms and no history of CHD but who were Need for Randomized Clinical Trial Data referred for exercise testing by their health care providers . 88 No increased risk of death associated with exercise-induced ischemia was found on echocardiography during a 10-year follow-up, even in the subgroups of individuals with several risk factors or who were at intermediate risk. By contrast, the DTS as well as resting left ventricular dysfunction (most of which was mild dysfunction) imparted increased risk. Thus, this study found no additional prognostic value to exercise echocardiography for screening of these relatively asymptomatic individuals but confirmed the value of exercise capacity or DTS in predicting risk.

By contrast, in a prior study of patients who were mildly symptomatic but at low pretest probability of disease, exerciseinduced ischemia on echocardiography was associated with increased risk of events during a 3-year period. 89 However, the event rate was low during follow-up, as was the sensitivity of the (47%), and the authors concluded that exercise echocardiography is not cost-effective in this setting.

There are several potentially important implications of the DIAD study for clinical guidelines and future design of clinical trials in asymptomatic individuals. First, the DIAD study demonstrated that it is feasible to conduct well-designed screening trials in asymptomatic individuals to determine the utility of screening tests with respect to clinical outcomes. Second, the low rates of cardiac events in asymptomatic individuals in DIAD, even when stress test results were abnormal, and the low positive predictive value of these abnormalities are consistent with results from prior prospective nonrandomized studies. Thus, it is not surprising that there was no clinical benefit associated with screening of these asymptomatic individuals, even though they were diabetic and had several other risk factors. Finally, this underscores the importance of conducting randomized clinical trials in asymptomatic populations to evaluate the clinical utility of screening stress testing, as called for by the AHA expert group. 5

CONCLUSION

Exercise stress testing is a safe procedure and provides both diagnostic and prognostic information in asymptomatic or 20. Diamond GA, Forrester JS: Analysis of probability as an aid in the clinical diagnosis of I coronaryminimally symptomatic individuals. A growing body of literature supports the utility of nonelectrocardiographic parameters—in particular, exercise capacity, chronotropic response, exercise blood pressure, and heart rate recovery—in determining prognosis in 22. Fearon WF, Lee DP, Froelicher VF: The effect of resting ST segment depression on the 29 asymptomatic subjects beyond traditional risk factors or global risk scores. However, data are still lacking that the screening use of 23. Rywik TM, O'Connor FC, Gittings NS, et al: Role of nondiagnostic exercise-induced ST-segment exercise testing results in improved patient outcomes. There are currently no Class I indications for exercise testing in asymptomatic adults in the ACC/AHA guidelines. Guidelines suggest that testing in asymptomatic individuals may be reasonable in the following situations: before vigorous exercise is started in the presence of diabetes, older age (men > 45 years, women > 55 years), or higher absolute risk for CHD due to comorbid conditions such as peripheral vascular disease and chronic kidney disease and for individuals involved in occupations potentially having an impact on public 27. Balady GJ, Larson MG, Vasan RS, et al: Usefulness of exercise testing in the prediction of safety. Exercise testing with imaging should be used as an initial test in the following situations : preexcitation syndrome (Wolff- 28. Gulati M, Arnsdorf MF, Shaw LJ, et al: Prognostic value of the Duke treadmill score in Parkinson-White), electrically paced ventricular rhythm, more than 1 mm of resting ST-segment depression, and complete left bundle 29. Rautaharju PM, Prineas RJ, Eifler WJ, et al: Prognostic value of exercise electrocardiogram branch block. Although some studies suggest additional prognostic value of imaging along with stress testing in asymptomatic subjects, 30. Ekelund LG, Haskell WL, Johnson JL, et al: Physical fitness as a predictor of cardiovascular stress testing with imaging is not cost-effective in this population because of consistently low event rates and low test specificity. Of note, the results of the DIAD trial further call into question the utility 31. Myers J, Prakash M, Froelicher V, et al. Exercise capacity and mortality among men referred of screening stress testing in asymptomatic diabetic patients. More data from randomized clinical trials in asymptomatic populations are needed to evaluate the utility of screening exercise testing.

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CHAPTER 30

Carotid Intima-Media Thickness Measurement and Plaque Detection for Cardiovascular Disease Risk Prediction

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KEY POINTS

- Measurement of carotid intimamedia thickness (CIMT) with Bmode ultrasound is a noninvasive, sensitive, and highly reproducible technique for identifying and quantifying arterial injury and cardiovascular disease risk.
- The relationship between increasing CIMT and incident cardiovascular disease events has been established across a wide age range; however, the strongest data are for individuals between 42 and 74 years of age.
- Several prospective, populationbased studies demonstrated that carotid plaque presence is associated with a significantly increased risk for myocardial infarction, stroke, and coronary heart disease death, independent of traditional risk factors.
- The relative risks associated with the presence of plaque are similar to or slightly higher than those observed with increased CIMT
- The most recent, comprehensive recommendations on the use of carotid ultrasound for cardiovascular disease risk assessment are contained in a Consensus Statement published in 2008 by the American Society of Echocardiography.
- The ASE Consensus Statement recommended that carotid ultrasound with CIMT measurement and evaluation for plaque presence be considered in intermediaterisk patients, in patients with a family history of premature cardiovascular disease in a first-degree relative, in individuals younger than 60 years old with severe

abnormalities of a single risk factor, and in women younger than 60 years old with two or more risk factors.

 Randomized outcome studies are needed to determine if improved risk prediction, behavior changes, and changes in physician practice that occur with CIMT or carotid plaque imaging lead to improved patient outcomes and cardiovascular disease risk reduction.

Cardiovascular disease (CVD) is the leading cause of death in the United States, accounting for approximately 864,480 (35.3%) deaths annually. 1 Each year, approximately 1.2 million individuals experience a myocardial infarction (MI), approximately one third of which is fatal. ¹Unfortunately, for a majority of individuals, the first symptom of heart disease sudden cardiac death or Atherosclerosis, the anatomical substrate for MI, begins in childhood and progresses over decades. 3 To prevent death and morbidity from coronary heart disease, there is great interest in identifying high-risk, asymptomatic patients who would be candidates for more intensive, evidence based medical interventions that prevent progression of atherosclerosis and reduce CVD risk. 4

Arterial imaging to identify and to quantify vascular disease has suggested to further refine coronary heart disease risk assessment. 4,5 As a screening test, imaging must be safe, sensitive, affordable, and lead to interventions that can favorably alter the natural history of CVD. Measurement of carotid intima-media thickness (CIMT) with B-mode ultrasound is a noninvasive, sensitive, and highly reproducible technique for identifying and quantifying arterial injury and CVD risk. It is a well-validated research tool that is increasingly being used as a clinical tool. 6-15 The United States Centers for Medicare and Medicaid has established a Current Procedural Terminology code (0126T) for "common carotid intima-media thickness (IMT) study for evaluation of atherosclerosis rhotic burden or coronary heart disease risk factor assessment.'

CAROTID INTIMA-MEDIA THICKNESS AND PLAQUE IMAGING

The extent of carotid artery atherosclerosis is strongly related to the extent of coronary artery atherosclerosis. 16 Carotid duplex ultrasound is used to evaluate for the presence and extent of occlusive, advanced, flowlimiting carotid artery atherosclerosis . 17 Ultrasound imaging of carotid artery wall thickness is a distinctly different test that is used to assess CVD risk. The superficial location and easy accessibility of the common carotid artery allow accurate, reproducible ultrasound imaging. The walls of the carotid artery are composed of the lumen-intima interface and the media adventitia interface that together produce two echogenic lines in the far wall of the carotid artery (Fig. 30-1). 9 Measurement of the combined thickness of these interfaces comprises the CIMT (Fig. 30-2). The presence of subclinical arterial injury is demonstrated by increased CIMT, and the presence of carotid plaque represents subclinical atherosclerosis disease.

RELATIONSHIP BETWEEN CAROTID INTIMA-MEDIA THICKNESS AND CARDIOVASCULAR DISEASE EVENTS

There are 11 published prospective, -population-based studies of CIMT and CVD risk that included at least 1000 participants and

FIGURE 30-1 Longitudinal plane demonstrating "double-line sign" on near and far walls of the common carotid artery.

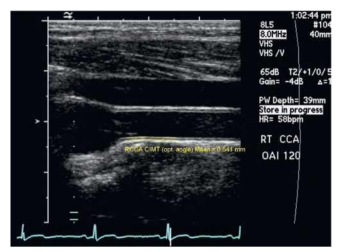


FIGURE 30-2 Leading edge-to-leading edge measurement of the far-wall common carotid artery CIMT.

presented odds ratios or relative risks adjusted for CVD risk factors (Table 30-1). 7,8,18-26 Many of these studies have recently been reviewed in detail. 27 These studies demonstrated that increased CIMT is associated with significantly increased risk for MI, stroke, and/or coronary heart disease death. 7,8,18 '26 Two additional large studies had similar findings. 28,29 In several studies, the adjusted relative risks associated with the greatest degrees of wall thickness (see cut points in Table 30-1) were high enough (> 2.0) that they would be expected to improve coronary heart disease risk prediction in appropriately selected patients. 7,8,18,20,23,29 Furthermore, CIMT me surements improve on traditional risk factors for classification of patients in regard to the presence of significant angio graphic coronary artery disease 30 and risk of recurrent CVD events. ³¹ In an analysis of unselected participants in the Atherosclerosis Risk in Communities (ARIC) study, CIMT values significantly (albeit modestly, ~0.02) increased the area under the receiver operator characteristic curve (AUC) for prediction of cardiovascular events in men. 32 In the Multi-Ethnic Study of Atherosclerosis (MESA), increased CIMT was positively associated with CVD events and stroke. 25 As in ARIC, CIMT values modestly increased the AUC (by ~0.01) for prediction of CVD compared with traditional risk factors alone. 25

The relationship between increasing CIMT and incident cardiovascular events has been established across a wide age range. However, the strongest data are for individuals between 42 and 74 years of age; several studies of individuals in this age range show similar results (see Table 30-1). For younger adults

(18 to 42 years old), consistent, strong relationships between increasing risk factor burden and CIMT, as well as between CIMT and emerging risk factors, have been demonstrated. ³³⁻⁴⁰ In the Carotid Atherosclerosis Progression Study (CAPS), CIMT predicted CVD events even among 2436 individuals < 50 years old (mean, 38.7 years; standard deviation, 7.0 years). ¹⁹ In that study, the relative risk associated with increased CIMT appeared to be higher among younger than older adults. ¹⁹

In the Tromso Study of 6226 subjects, there were conflicting data regarding the predictive value of CIMT and CVD events. The CIMT in the common carotid artery did not demonstrate a statistically significant relationship between CIMT and first MI, ²⁴ but there was a significant relationship, in both men and women, when CIMT was evaluated with the use of both the common carotid artery and bulb segments. 24 However, this study measured CIMT only in the right carotid artery. The American Society of Echocardiography (ASE) Consensus Statement recommends that CIMT be evaluated with the far wall of both the right and left common carotid arteries, in conjunction with plaque imaging in the common carotid, bulb, and internal carotid artery segments, because atherosclerosis progresses more rapidly in the bulb and internal carotid artery segments, and the effects of risk factors on CIMT can vary by segment. 9,41-44 This approach has been validated, and its predictive value has been rigorously demonstrated in other studies (see Table 30-1). 9

RELATIONSHIP BETWEEN CAROTID PLAQUES AND CARDIOVASCULAR DISEASE EVENTS

Carotid plaque presence is associated with the presence of coronary artery plaque and the occurrence of future cardio vascular and cerebrovascular disease events. 24,26,30 Most carotid plaques are found in the bulb and proximal internal carotid artery segments because of turbulent flow. 45 Atherosclerosis and CIMT progress more rapidly in the bulb and internal carotid segments compared with the common carotid artery. 41,42 Seven population-based studies that included at least 1000 participants and presented relative risks or hazard ratios adjusted for CVD risk factors have demonstrated the predictive power of carotid plaque (Table 30-2). 21-23,26,46-48 In these studies, the relative risks associated with the presence of plaque were similar to or slightly higher than those observed with increased CIMT. Two additional large studies of carotid plaque presence and two studies that evaluated plaque area had similar results. 24,29,49,50 The presence of carotid plaque predicted future CVD events among young, middle, and elderly subjects. ^{24,49} A prospective study of 367 elderly men (> 70 years old) demonstrated that the presence of carotid plaque significantly improved the AUC for prediction of all cause mortality and cardiovascular mortality, even after considering traditional cardiovascular risk factors and use of medications (by ~0.03 to 0.04). ⁴⁹ In San Daniele Township, the presence of carotid plaque or CIMT > 1 mm had greater predictive value than the Framingham risk score for ischemic cerebrovascular disease events and a small increase in AUC (by ~ 0.01). ²⁶

Unfortunately, the definition of carotid plaque in most reported studies was not uniform. ⁵¹ Most studies identified plaque as focal widening relative to adjacent segments with protrusion into the lumen and/or had a minimum wall thickness. ⁵¹ The Mannheim Carotid Intima-Media Thickness Con sensus Report suggested that plaque should be defined as "a focal structure that encroaches into the arterial lumen of at least 0.5 mm or 50% of the surrounding IMT or demonstrates

	Disea	se (N / 10	000 participants	each)			
Study ARIC 7	N 12,841	Age, yea (% W) 45-64 (57%)	rs Follow-up, years 5.2	Measurement and Site(s) Mean of mean CCA, bulb, ICA	Eventos MI, CHD death	Adjusted RR (95% CI)	CIMT Cut Point, Adjusted RR (95% CI) ·* Highest tertile W: 2.53 (1.02-6.26) M: 2.02 (1.32-3.09)
				Mean Approx	MI, CHD death	0.19 W: 1.38 (1.21-1.58) M: 1.17 (1.04-1.31) 0.19 W: 1.46 (1.22-1.74) M: 1.08 (0.91-1.1.27)	
ARIC 18	14,214	45-64 (55%)	7.2	Mean of mean CCA, bulb, ICA	Stroke	0.19 W: 1.36	Highest tertile W: 2.32 (1.09-4.94) M: 2.24 (1.26-4.00)
				Mean Approx	Stroke	(1.16-1.59) M: 1.21	Highest tertile W: 1.65 (0.85-3.19) M: 2.69 (1.49-4.87)
CHAPTER 19	5056	19-90 (49.7%)	4.2	Mean Far-wall CCA Mean Far-wall CCA Mean Far-wall CCA	ml Stroke Ml, stroke, death	0.16 1.16 (1.05-1.27) 0.16 1.11 (0.97-1.28) 0.16 1.17 (1.08-1.26)	Highest quartile 1.83 (0.97-3.45) Highest quartile 1.82 (0.64-5.16) Highest quartile 1.85 (1.09-3.15)
CHS :	4476	> 65 (39%)	6.2	Mean of maximum Near + far CCA, ICA Maximum Near + far CCA Mean of maximum Near + far CCA, ICA Maximum Near + far CCA	ml Stroke Stroke	1 SD 1.36 (1.23-1.52) 0.20 1.24 (1.12-1.38) 1 SD 1.33 (1.20-1.47) 0.20 1.28 (1.16-1.42)	Highest quintile 3.61(2.13-6.11) Highest quintile 2.46 (1.51-4.01) Highest quintile 2.57 (1.64-4.02) Highest quintile 2.13 (1.38-3.28)
(IHD ∞	1257	42-60 (0%)	3	Maximum Far-wall CCA	ml	0.11 1.11 (1.06-1.16)	> 1.0mm 2.1 (0.8-5.2)
ADCS 22	5163	46-68 (60%)	7	Maximum Far-wall CCA	MI, CHD death	0.15 1.23 (1.07-1.41)	Highest tertile 1.50 (0.81-2.59)
ABLE 25	6698	48-84	5.3	Mean of maximum Near + far CCA, ICA	CHD, stroke, cardiovascular death	1 SD 1.2 (1.0-1.3)	Highest quartile 1.7 (1.2-2.5)
Rotterdam 23	6389	> 55 (62%)	7-10	Maximum Near + far CCA	ml	0.21 1.28 (1.14-1.44)	Highest quartile 1.95 (1.19-3.19)
San Daniele 28	1348	18-99 (53%)	12.7	Mean of maximum Far-wall CCA	Ischemic stroke, TIA, vascular death	-	> 1.0mm 5.6 (3.2-10.1)
romsø 24	6226	25-84 (52%)	5.4	Mean of mean Near + headlight CCA, bulb, ICA	ml	_	Highest quartile W: 2.86 (1.07-7.65) M: 1.73 (0.98-3.06)
Yao City 21	1289	60-74 (0%)	4.5	Mean of maximum Near + far CCA, ICA	Stroke	-	Highest quartile 4.9 (1.9-12.0)

TABLE 30—1 Prospective Studies of Carotid Intima-Media Thickness and Risk for Cardiovascular Disease Events in Individuals Without Known Cardiovascular

a thickness of > 1.5 mm." 52,53 An ASE report defined nonobstructive plaque as "the presence of focal thickening at least 50% greater than that of the surrounding vessel wall." 54 These definitions are similar to those used in the ARIC study, the largest prospective cohort study that demonstrated the predictive value of plaque in cardiovascular risk assessment 46

The ASE Consensus Statement recommended that carotid plaque be defined as the presence of focal wall thickening that is at least 50% greater than that of the surrounding vessel wall or as a focal region with CIMT > 1.5 mm that protrudes into the lumen and is distinct from the adjacent boundary. 9,46,52-54

Adjusted for age, sex, and traditional risk factors.

[^] Highest tertile or quartile compared with lowest tertile or quartile.

CCA, common carotid artery; CHD, coronary heart disease; CI, confidence interval; CIMT, carotid intima-media thickness; ICA, internal carotid artery; M, male; MI, myocardial infarction; RR, relative risk; SD, standard deviation; TIA, transient ischemic attack; W, women.

		Age, years	(70 I Onow up,		Flaque Fresence, Aujusteu	riaque rresence, Aujusted
Study	N	W)	years	Eventos	HR (95% CI) ·	RR (95% CI) ·
ARIC 46	12,375		7	MI, CHD death		_
		45-64 (54%)			With AS: 2.96 (1.54-3.30) Without AS: 2.02 (1.42-2.41)	
KIHD 47	1288		< 2	ml	4.15 (1.5-11.47)	_
		42-60 (0%)				
MDCS 22	5163		7	MI, CHD death	1.81 (1.14-2.87)	_
		46-68 (60%)				
Northern Manhattan 48	1939		6.2	Stroke	3.1 (1.1-8.5)	_
		> 40 (59%)				
Rotterdam 23	6389		7-10	ml		_
		> 55 (62%)			severity 1.83 (1.27-2.62)	
San Daniele 28	1348	18-99 (53.3%)	12.7 (average)	Ischemic stroke, TIA, vascular death	-	10.4 (6.4-17.1)
Yao City 21	1289	60-74 (0%)	4.5	Stroke	-	3.2 (1.4-7.1)

· Adjusted for age, sex, and traditional risk factors.

AS, acoustic shadowing; CHD, coronary heart disease; CV, cardiovascular; HR, hazard ratio; RR, relative risk; MI, myocardial infarction; TIA, transient ischemic attack; W,

Quantification of carotid plaque area is also a strong predictor of future CVD events. 24,50,55 In some studies, the two dimensional total carotid plaque area was more predictive of MI than CIMT, after adjustment for traditional CVD risk factors. 24,56 In the Tromso Study, baseline carotid plaque area was a stronger predictor of future MI than CIMT (see Tables 30-1 and 30-2), especially in women. In both men and women, there was a direct correlation between total plaque area and MI incidence. 24 In a report of 1686 individuals in an atherosclerosis prevention clinic, those in the highest quartile of carotid plaque had the greatest risk of future vascular events (see Table 30-2), and subjects with progression of carotid plaque area doubled their risk of future events compared with subjects with stable plaque area. 24

Disease (N > 1000 participants each)

The ASE does not currently recommend risk stratification based on carotid plaque measurements outside of the research setting because of their complex geometry and the absence of a published, widely applicable standard for imaging and measurement of carotid plaque area. 54,57 Carotid plaque area and even volume have been used in limited clinical settings to assist with risk stratification and to evaluate the efficacy of atherosclerosis disease management. 12,50,55,58 However, the incremental predictive value of the quantification of carotid plaque, beyond defining the presence of plaque, is unknown.

GUIDELINE AND CONSENSUS STATEMENT RECOMMENDATIONS FOR CAROTID INTIMA-**MEDIA THICKNESS**

The use of carotid ultrasound for evaluation of CIMT and carotid plaque as a clinical risk prediction tool has been addressed in several guidelines and consensus statements. 9 In 2000, the American Heart Association Prevention Conference V concluded that CIMT "can now be considered for further clarification of coronary heart disease risk assessment at the request of a physician," provided it is performed by an experienced laboratory. 4 In 2001, the National Cholesterol Education Program Adult Treatment Panel III stated that CIMT "could be used as an adjunct in coronary heart disease risk assessment ... the finding of an elevated carotid IMT (eg, > 75th percentile for age and sex) could elevate a person with multiple risk factors to a higher risk category." 59 This expert panel concluded that "if carried out under

proper conditions, carotid IMT could be used to identify persons at higher risk than that revealed by the major risk factors alone." ⁵⁹ In 2003, the 34th Bethesda Conference supported the use of CIMT as a screening test for subclinical vascular disease. ⁵ The clinical application of CIMT methodology was reviewed and supported in 2006 in a report from the American Society of Echocardiography and the Society of Vascular Medicine and Biology. 54 In 2007, the European Society of Cardiology described increased CIMT as a marker of hypertensive target organ damage. 60 The most recent, comprehensive recommendations on the use of carotid ultrasound, including CIMT and carotid plaque presence, for cardiovascular risk assessment are contained in a Consensus Statement published in 2008 by the American Society of Echocardiography and were endorsed by the Society of Vascular Medicine. 9

APPLICATION OF CAROTID ULTRASOUND FOR CARDIOVASCULAR DISEASE RISK **ASSESSMENT**

Carotid ultrasound is a noninvasive tool to identify asymptomatic patients at increased CVD risk. In traditional CVD risk assessment, risk factors are used to estimate a 10-year CVD event risk; however, this estimate may not accurately reflect elevated long-term risks, especially in young and middle-aged adults, women, and ethnic minorities. 61-68 A longitudinal study that included subjects from the Coronary Artery Risk Development in Young Adults (CARDIA) and MESA studies demonstrated that subjects with a low 10-year but high lifetime CVD risk had higher common and internal CIMT compared with those at low lifetime risk and had greater progression of coronary artery calcification during 15 years. 69 Cross-sectional and longitudinal studies have integrated CIMT with CVD risk algorithms and assessment demonstrated reclassification 10-13 and improvement in CVD risk stratification by individualizing vascular risks with use of CIMT results, especially in intermediate-risk patients. 6,14,70

The ASE Consensus Statement 9 recommended that carotid ultrasound with CIMT measurement and evaluation for plaque presence could be considered in the following types of patients if the level of aggressiveness of therapy (eg, phar macotherapy) or additional information about the burden of subclinical vascular disease or future CVD risk is needed:

- · intermediate risk (Framingham risk score 6% to 20% without established coronary heart disease, peripheral arterial disease, cerebrovascular disease, diabetes mellitus, or abdominal aortic
- family history of premature CVD in a first-degree relative (men < 55 years old, women < 65 years old);
- individuals < 60 years old with severe abnormalities in a single risk factor (such as genetic dyslipidemia) who are not being treated with medications; hours
- women < 60 years old with at least two CVD risk factors.

COMPARISON OF CAROTID INTIMATE MEDIA THICKNESS AND CORONARY ARTERY **CALCIUM**

Fast computed tomography for measurement of coronary artery calcium is another technique that evaluates for the presence and extent of subclinical vascular disease and predicts future CVD events. 71-73 A report from the MESA study compared the prognostic value of CIMT and coronary artery calcium in predicting fatal CVD events, development of coronary heart disease, and stroke in 6698 participants (45 to 84 years old) during 5.3 years of follow-up. 25,74 Both imaging modalities predicted the primary outcomes; however, coronary artery calcium was a stronger predictor of incident CVD than was CIMT. After adjustment for each other (coronary calcium and CIMT) and traditional risk factors, the hazard ratio for each standard deviation increase in coronary artery calcium increased by 2.1fold compared with 1.3-fold for each standard deviation increase in maximum CIMT, and the change in the AUC for prediction of CVD events was greater with coronary artery calcification (\sim 0.03) than with CIMT (~0.01). 25 CIMT, but not coronary artery calcification, predicted incident stroke. 25 In the Cardiovascular Health Study (CHS) of adults older than 65 years old, common carotid artery CIMT (hazard ratio, 11.25 [2.28-55.61]) was a stronger predictor of strokes compared with coronary artery calcium (hazard ratio, 3.73 [0.81-17.11]). 75 The hazard ratios for coronary heart disease and total CVD were not statistically different between CIMT and coronary artery calcium.

The clinical predictive value of coronary calcium in younger men, women, and African Americans is less clear because of a lower prevalence of coronary artery calcium in these subgroups. CIMT has the advantage of being a continuous measure that could be used, if clinically indicated, to stratify risk in individuals for whom coronary artery calcium scoring may have limited discriminatory power because of a high predicted prevalence of a zero calcium score. ⁷⁶ In a report from MESA, increased CIMT predicted total CVD events (hazard ratio, 2.2; 95% confidence interval, 1.2-4.0) among individuals with low coronary artery calcium scores (< 10), along with increasing age, family history of CVD, use of lipid-lowering medications, and smoking. 77 CIMT (hazard ratio, 3.2; 95% confidence interval, 1.5-6.6) and former and current smoking were independent predictors of hard CVD events. 77

As a screening test, carotid ultrasound has additional advantages that should be considered. Although the amount of radiation exposure with modern calcium screening by noncontrast-enhanced fast computed tomography is relatively low, carotid ultrasound does not involve any exposure to ionizi568 greater perceived likelihood of having or developing heart disease

young and middle-aged adults. 78 Compared 507 with computed tomography, CIMT is an inexpensive, portable predictive tool that can be used in the office setting. 79 Also, carotid ultrasound can be performed serially to monitor the clinical effectiveness of therapeutic intervention or changes in arterial injury over time without risk of recurrent radiation exposure 55; however, serial monitoring is not recommended by the ASE Consensus Statement at this time. 9

EFFECTS OF CAROTID INTIMA-MEDIA THICKNESS SCREENING PROGRAMS ON **CLINICAL PRACTICE AND OUTCOMES**



Several clinical CVD risk assessment programs have used carotid 30 ultrasound to measure CIMT. 6,10-15,80 In clinical practice, CIMT has been shown to help reclassify patients at intermediate risk, ^{6,10-12} to discriminate between patients with and without prevalent CVD, ²⁸ and to predict major adverse cardiovascular events. ^{6,14} Most of these studies considered the patient's age and sex by using normative percentile values. 10,11,13-15 The METEOR (Measuring Effects on Intima Media Thickness: An Evaluation of Rosuvastatin) study demonstrated that middle-aged adults at apparently low to intermediate CVD risk but with increased CIMT benefited from statin therapy that they otherwise would not have qualified for on the basis of current treatment guidelines. ^{59,81} In this prospective, randomized multicenter clinical trial, the magnitude of the difference in CIMT progression rates (-0.145 mm/year) was similar to that observed in secondary prevention trials that were associated with a reduction in CVD events. 81,82 Although not definitive, this study suggests that use of CIMT to modify preventive treatment strategies is feasible and associated with a delay in the progression of arterial wall thickening, which is associated with reduced CVD risk.

Some data suggest that increased CIMT or carotid plaque presence can influence the behavior of patients and physicians. A small study (N = 153) randomly assigned smokers to a smoking cessation intervention or smoking cessation plus carotid ultrasound, after which they were shown a picture of their carotid plaque. Smoking cessation rates were 22.2% in the 54 participants with at least one plaque compared with 5% and 6.3% among those without plaque or who did not have ultrasonography (P = 0.003). 83 In a small (N = 23), pilot study of visual feedback from carotid ultrasound imaging compared with verbal feedback, the intervention increased perception of smoking-related illness and smoking cessation behavior and intention; however, intention increased only in people with high levels of self-efficacy (P < 0.03). 84 In a study of 210 individuals reported in abstract form and described in a review paper, patients were more likely to adhere to recommendations for diet, exercise, and smoking cessation 12 months after seeing pictures of their CIMT examination. 85

In another promising pilot study, carotid plaque screening was performed on asymptomatic patients with at least two CVD risk factors to assess the impact of carotid plaque screening on the physician's and patient's behavior in an office practice setting. Identification of carotid plaques increased the likelihood that physicians would prescribe aspirin (P = 0.031) and lipid-lowering therapy (P = 0.004), but did not change patients' motivation to make lifestyle changes. 86 In a subsequent multicenter trial of 253 patients, CIMT and carotid plaques were assessed in a clinical office setting by non-sonographer clinicians using a handheld ultrasound system. When increased CIMT or carotid plaque was detected, physicians significantly changed their behavior by ordering aspirin and lipid-lowering therapy (odds ratios, 2.9-7.4; P < 0.001), interventions that are proven to reduce CVD risk in patients at increased risk. ⁷⁹ Furthermore, patients had a

radiation, an important consideration when imaging healthy = 0.004) and reported greater intentions to take cholesterol-lowering

medication (P=0.002); however, even subjects without ultrasound abnormalities reported increased motivation to exercise (P=0.003) and make dietary changes (P=0.051), illustrating the complexity of affecting a patient's behavior with imaging. ⁷⁹ The coronary artery calcium screen ing literature also suggests that the behavior of physicians is affected more than that of patients and is missing long-term data showing that a strategy for atherosclerosis screening is superior to current strategies or case management. ⁸⁷⁻⁹¹ Large, randomized outcome studies are needed to determine if improved risk prediction, behavior changes, and changes in the practice of physicians that occur with CIMT and/or 30 carotid plaque imaging translate into improved patient outcomes and reduced cardiovascular events.

CAROTID INTIMA-MEDIA THICKNESS AND CAROTID PLAQUE SCANNING TECHNIQUE

The following recommendations are from the recent ASE Consensus Statement. ⁹ Before the study is started, both the sonographer and patient must be positioned comfortably to facilitate accurate, high-quality, and reproducible images. The patient should lie supine with their head on the scan table. The ideal position of the neck is a slight hyperextension and rotation in the direction opposite the test. Use of external landmarks and a 45-degree angle wedge pillow can help standardize the transducer's angle relative to the patient's head position (Fig. 30-3). Standard three-lead electrocardio graphic monitoring should be performed with acquisition of easily discernible R-wave deflections. Scan time can vary according to protocol requirements and sonographer experience. ⁹²

Carotid arteries should be imaged with a state-of-the-art ultrasound system with a linear array transducer operating at a fundamental frequency of at least 7 MHz. Most subjects can



FIGURE 30-3 Position of the patient and setup for carotid ultrasound study. be scanned at a standard depth of 4 cm from anterio.

be scanned at a standard depth of 4 cm from anterior to posterior planes. With increased depth, there is decreased resolution - (larger pixel size). A single focal zone, high dynamic range, and fundamental frequencies without harmonic or compound imaging should be used. Use of ultrasound contrast, although a promising research technique, is not recommended for clinical assessment of CIMT at this time. B-mode imaging, rather than M-

mode imaging, is recommended. All reported observational studies relating CIMT values to car diovascular events used B-mode measurements averaged over at least a 1-cm segment. Multiple measurements of multiple extended segment lengths allow expression of CIMT values at higher levels of precision instead of simple multiples of pixel size.

Carotid ulfrasound imaging should follow a protocol from a large epidemiological study that reported CIMT values in percentiles by age, sex, and race or ethnicity. ARIC, MESA, Bogalusa Heart Study, and CHS are large, cross-sectional, highquality studies that reported common carotid artery CIMT values by age, sex, and race or ethnicity and were conducted in North America. 8,9,41,93 The 50th percentile mean far-wall common carotid artery CIMT values for white and black men and women between 45 and 64 years of age in ARIC and MESA were quite similar, given the differences between the studies. 9,41 For older patients, the 50th percentile maximum common carotid artery CIMT values in the CHS tended to be higher than in MESA, probably because individuals with known CVD were excluded from MESA but not from CHS. 8,9 Several large, high-quality studies from Europe also reported common carotid artery CIMT values. 9,19,22,94 These studies did not provide information about race or ethnicity but were conducted mostly in white individuals. In general, CIMT values in studies such as CAPS and the Malmo Diet and Cancer Study (MDCS) tend to be higher than in the North American studies. 9,19,22 Reasons for the thicker CIMT values observed in these studies include different population characteristics, instrumentation, imaging and recording standards, and measurement techniques, such as whether segments with plaque were included or excluded from the measurement protocol. The relative risks associated with increasing CIMT were similar across all of the studies in Table 30-

The ASE Consensus Statement recommends that CIMT of the distal 1 cm of the far wall of each common carotid artery be imaged and compared with a normative data set. 9 The distal common carotid artery is easy to image, and far-wall common carotid artery CIMT measurements predict future cardiovascular events (see Table 30-1). Although near-wall measurements and those from other segments have also been used in some studies, they are more challenging technically, less reproducible, and do not appreciably improve risk prediction. 54 In one study, the ultrasound measurement of the near-wall CIMT was 20% lower than the corresponding histological measurement. 95 Although CIMT imaging is limited to the common carotid artery, a thorough scan of the extracranial carotid arteries for the presence of carotid plaque should be performed to increase sensitivity for identification of sub-clinical vascular disease. A circumferential plague scan of both carotid arteries can compensate for reduced sensitivity that may result from measurement of only common carotid artery CIMT. 54,96

A scanning protocol based on that used in the ARIC study was recommended by the ASE Consensus Statement because in ARIC, prevalent common carotid artery CIMT and carotid plaques predicted future cardiovascular events (see Tables 30-1 and 30-2), and the scanning methods are reproducible in most clinical laboratories. ^{7,9,18,41,46,97} Furthermore, ARIC was a very large study, and normative values based on age, sex, and race or ethnicity have been published in the age range that is considered most appropriate for screening. ⁴¹ The

recommended protocol includes a plaque scan and imaging of the common carotid artery from multiple angles for measurement of CIMT. The ARIC scanning protocol, limited to the common carotid artery, is summarized in the ASE Con sensus Statement. 9 Additional information about frequently observed pitfalls and possible solutions to CIMT acquisition problems are provided in that document. Other scanning protocols may be used if they are more germane to the clinical population being investigated. Which segments of the carotid artery are interrogated and at which angles, as well as which measurements are obtained, must match those in the normative data set of the representative epidemiological study. The age, sex, and race or ethnicity of the patient should be considered in choosing the scanning and measuring protocols. For example, the CIMT measurements in the Bogalusa Heart Study have not yet been related to future CVD events; however, they are the only normative values for CIMT in young adults from North America. 93 The MESA study has the only normative values for CIMT in Chinese and Hispanic Americans. On the basis of the relatively similar relative risk associated with increasing CIMT across the age ranges described in Table 30-1, it can be inferred that increased CIMT in these patient groups (as determined by comparison with these data sets) is associated with increased cardiovascular risk. The use of values from clinically referred populations is discouraged because of the likelihood of referral bias and inaccurate risk estimates.

CAROTID INTIMA-MEDIA THICKNESS MEASUREMENT

The ASE Consensus Statement recommends the use of digital images that are acquired and stored directly from the ultrasound system, rather than digitized video captures, as well as minimal compression and use of the DICOM standard. 9 The reader reviews overall image quality, thickness, presence or absence of plaque, and presence or absence of incidental findings such as possible obstructive carotid artery disease, carotid dissection, carotid tumors, thyroid abnormalities, and lymphadenopathy. Cine loops and R wave-gated still-frame images of the distal 1-cm segment of the far walls of the right and left common carotid artery, each from three different angles of incidence, are measured in triplicate. Measurement of each image involves tracing of the blood-intima and media adventitia interfaces of the far wall by a leading edge-to-leading edge method (see Fig. 30-2). ⁹Only clearly visualized images should be measured. If plaques are detected in the segment being measured, they are traced as part of the CIMT because they appear to have been included in CIMT measurements in most of the epidemiological studies in Table 30-1. 27 Most studies that provided reference values were obtained bv manual reading techniques: however. semiautomated border detection programs were used by some. 9 Semiautomated border detection programs are widely available, and when they are used, they tend to improve reproducibility and to shorten reading time, especially among newer readers. 9,98 101 The ASE Consensus Statement recommends use of a semiau tomatod border detection program with validated accuracy. 9 These programs tend to produce somewhat thicker CIMT values than are seen with manual tracing, especially if the generated borders are left unedited. Simple point-to-point measurement of CIMT should be avoided.

Depending on the protocol, the triplicate measurements are averaged or the highest value is taken. Most reading software will report the mean-mean (average of segmental mean CIMT values) and mean-maximum (average of segmental maximum CIMT values) thickness values. Mean-mean values are more reproducible because multiple points along the traced segment are averaged, but they are less sensitive **509** to change. Mean-maximum values are more sensitive to change but less

reproducible because they are derived from a single-point measurement along the 1-cm region. It is recommended that mean CIMT values from the far walls of the right and left common carotid arteries (mean-mean) be used for clinical studies.

STUDY INTERPRETATION

The study report (1) provides the referring health care provider with data about the mean (or maximum) CIMT of the segments analyzed, depending on the scanning protocol and reference database; (2) briefly summarizes the scanning protocol and reference data base used; (3) states whether the CIMT values are means or maxima; (4) compares the CIMT data with the reference population of individuals of similar age, sex, and race or ethnicity; (5) describes the presence or absence of carotid plaques; and (6) describes any other clinically relevant findings, such as the presence of obstructive carotid atherosclerosis. Each of these points is discussed in detail in the ASE Consensus Statement. ⁹To avoid confusion with a duplex carotid ultrasound examination, the report should clearly identify the type of study being performed (ie, "carotid ultrasound study for cardiovascular risk assessment ") and explicitly state that the study measures the thickness of the walls of the carotid artery and that the results are not a "percent stenosis" and do not indicate the presence or absence of clinically significant obstruction, unless noted otherwise. 9 Reports should also include a clinically relevant percentile range for the CIMT value by stating a range of percentiles (such as quartile or quintile) for the measured value. 9 The communication of CIMT results can be facilitated by qualitatively describing broad ranges of percentiles to avoid the appearance of greater precision that is achievable when CIMT values are mapped to a reference population. This is because the percentile estimates in the population studies have confidence intervals surrounding them and because the instrumentation, scanning, and measurement techniques in a clinical laboratory will not be exactly the same as those used in these studies. 9 Values > 75th percentile are considered high and indicative of increased risk. Values in the > 25th percentile but < 75th percentile are considered average and indicative of unchanged risk. 9 Values < 25th percentile are considered low, but whether they are indicative of reduced risk is unclear.

CONCLUSION

Evaluation of CIMT and carotid plaque by B-mode ultrasound is a validated tool for CVD risk assessment that can be integrated into clinical practice. This noninvasive approach can detect subclinical vascular disease and refine CVD risk assessment in some asymptomatic patients. Strict attention to quality control in image acquisition, measurement, interpretation , and reporting is necessary for the implementation of this technique in clinical practice.

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CHAPTER 31

Peripheral Arterial Disease Assessment and Management

Jeffrey S. Berger and Emile R. Mohler III

KEY POINTS

- Lower extremity peripheral arterial disease (PAD) is a highly prevalent condition in the United States.
- The risk factors for PAD are similar to those for arterial disease in the coronary and cerebrovascular territories.
- There is growing evidence that several nontraditional risk factors, including inflammation, endothelial dysfunction, thrombotic and hemostatic markers, and platelet activity, may also be associated with increased PAD risk.
- After a complete history and physical examination, the diagnosis of PAD should be confirmed with a vascular study.
- The treatment of PAD includes a broad approach, focusing on reduction of the risk of the major factors associated with the development and progression of PAD.
- The treatment of lower extremity, stable, claudication symptoms from PAD involves an initial approach of exercise and pharmacologic therapy and, if it is not successful, selective revascularization based on anatomy and lifestyle considerations.
- Patients with critical limb ischemia should undergo immediate evaluation for possible revascularization.
- The evaluation for an abdominal aortic aneurysm involves physical examination and abdominal ultrasonography.

Peripheral arterial disease (PAD) is a disease that obstructs blood supply to the extremities . The most common pathological process is atherosclerotic disease, but it may also result from connective tissue diseases, vasculitis, hematological disorders, thrombo fibromuscular dysplasia, embolism, mechanical obstruction, and occupationrelated diseases. This serious and highly prevalent disorder is associated with significant morbidity and mortality, commonly with impaired function and quality of life.

EPIDEMIOLOGY

Lower extremity PAD, a highly prevalent condition in the United States with an estimated general population prevalence of 12%, is estimated to affect approximately 8 to 12 million Americans. ¹ In primary care office practices, PAD is present in 29% of patients older than 70 years or older than 50 years with a history of smoking or diabetes . 2 The prevalence of PAD and intermittent claudication increases progressively with age. Data from the 1999-2000 National Health and Nutrition Examination Survey (NHANES) demonstrated the prevalence of PAD (defined as an ankle-brachial index < 0.90 in either leg) was 0.9% between the ages of 40 and 49 years, 2.5% between the ages of 50 and 59 years, 4.7% between the ages of 60 and 69 years, and 14.5% at the age of 70 years and older. 3 NHANES data from 1999 to 2004 demonstrated a 44% increase in the odds of PAD for every 10-year increase in age. ⁴ There are data to suggest that race or ethnicity plays a role in PAD. 3,5-7 The prevalence among non -Hispanic blacks was 7.9%, the highest among other racial and ethnic groups (4.4% in non-Hispanic whites and 3% in Mexican Americans). Consistently, the Atherosclerosis Risk in Communities (ARIC) study demonstrated a significantly higher prevalence of PAD in African Americans than in whites. 8 Collaborative data from seven community-based studies noted a higher rate of PAD in African Americans than in American Indians, Asian Americans,

Hispanics, and non-Hispanic whites. 9

Despite the high prevalence of PAD, detection and awareness are lower than for arterial disease in other locations. ² A population-based telephone survey of a nationally representative sample of adults older than 50 years demonstrated that only 26% of respondents expressed familiarity with PAD. ¹⁰

RISK FACTORS FOR PERIPHERAL ARTERIAL DISEASE

The risk factors for PAD are similar to those for arterial disease in the coronary and cerebrovascular territories. 5,11 Increasing age, hypertension, dyslipidemia, cigarette smoking, and diabetes mellitus are wellestablished risk factors for all arterial disease. Data derived from several observational studies demonstrated that cigarette smoking and diabetes are particularly strong risk factors for PAD. Data from the 1999 2000 NHANES survey found that in age- and sexadjusted logistic regression analysis, current smoking had a greater than fourfold increased odds and diabetes almost a threefold increased odds of prevalent PAD. In a longitudinal cohort study, data from the Framingham Heart Study of 381 men and women who were observed for 38 years revealed that the odds ratio for development of intermittent claudication was 2.6 for the presence of diabetes mellitus and 1.4 for each 10 cigarettes smoked per day. 12

Dyslipidemia is also associated with an increased prevalence of PAD. ⁵ In comparison of a panel of lipid risk factors for PAD, total cholesterol, high-density lipoprotein cholesterol (HDL-C), low-density lipoprotein cholesterol (LDL-C), triglycerides, and apolipoprotein B were all significant predictors of increased risk of PAD, although the ratio of total cholesterol to HDL-C was the single strongest predictor (highest quartile versus lowest quartile relative risk, 3.9; 95% confidence interval, 1.7 to 8.6). ¹³

inflammation, endothelial dysfunction, development and eventual rupture. 34 including thrombotic and hemostatic markers, and platelet activity, may also PAD.

In the Edinburgh Artery Study, 16 a population-based cohort mechanisms and PAD. study, several markers of endothelial function and inflammation were compared. C-reactive protein, interleukin-6, and intercellular muscle structure and function. Repeated episodes of ischemia adhesion molecule 1 were significant predictors of lower extremity during exercise and reperfusion during recovery may | promote atherosclerotic progression measured by ankle-brachial index oxidant injury to endothelial cells, muscle mitochondria, muscle during 12 years of follow-up independently of cardiovascular risk fibers, and distal motor axons. Muscle denervation and alterations factors. Elevated levels of inflammatory biomarkers are also in muscle metabolism contribute to performance limitations. 38 associated with greater functional impairment and faster functional decline in people with PAD. 17,18 A study of proteomic profiling limb ischemia. Patients with claudication frequently progress to identified the protein § 2-thromboglobulin as being elevated in chronic limb ischemia and have a combined annual amputation subjects with PAD, and a significant correlation was observed mortality rate of 2% to 4% per patient per year, whereas patients between \$ 2 - microglobulin and the severity of disease. 19 The with chronic limb ischemia have a 6-month amputation risk of association of p₂-microglobulin with PAD may be related to 25% to 40% and an annual mortality rate as high as 20% (Fig. 31vascular inflammation.

Abnormalities in hemostasis also associate with an increased prevalence of PAD. Data derived from the ARIC study found small-vessel PAD. In a longitudinal study of 403 subjects with a elevated levels of hemostatic markers in subjects with PAD. mean follow-up of 4.6 years, current cigarette smoking, the ratio Specifically, higher levels of fibrinogen, von Wil lebrand factor, of total cholesterol to HDL-C, high-sensitivity C-reactive protein, factor VIII, D-dimer, and thromboglobulin were associated with and lipoprotein(a) were independent predictors of large-vessel greater PAD prevalence. In a population of Scottish men and PAD progression. This was in contrast to progression of smallwomen aged 55 to 74 years, mean levels of fibrinogen, fibrin D- vessel PAD, for which diabetes was the only significant dimer, and plasma viscosity remained significantly higher among predictor. 39 the diabetes/impaired glucose tolerance group with PAD compared with those with no PAD. 20

In contrast, fibrinogen and homocysteine were not associated DIAGNOSIS OF PERIPHERAL with the development of PAD in the Women's Health Study. ARTERIAL DISEASE Considerable evidence links platelets, a major culprit in atherothrombosis, and the development of PAD. With use of data After a complete history and physical examination, the diagnosis from NHANES, levels of mean platelet volume in peripheral blood of PAD should be confirmed with a vascular study. There are independently associated with PAD (tertile 1, 4.4%; tertile 2, 6.1%; multiple modalities from which to choose for assessment of PAD. tertile 3, 7.0%; P for trend = 0.003), independent of traditional A simple, inexpensive, noninvasive tool that correlates well with cardiovascular risk factors. 21 Ongoing studies will determine the angiographic disease severity and functional symptoms is the usefulness of markers of inflammation, endothelial function, hemostasis, and platelet activity in the risk prediction of PAD.

PATHOPHYSIOLOGY

Although it is poorly understood, the pathophysiological process leading to development of claudication and decline in functional status is thought to be progression of athero-thrombosis, ^{22,23} the of atherosclerosis and throm bosis. be classified into proatherogenic state can increased vasoconstriction, inflammation, and platelet activation and 40,45,46 ABI values between 0.90 and 0.99 are considered borderline endothelial dysfunction. The natural history of atherosclerosis in and equivocal for PAD. Of note, recent data indicate that even the extremities involves progressive occlusion of the vessel, borderline ABI values have significantly increased risk for a typically in susceptible regions where turbulent blood flow occurs, cardiovascular event (Fig. 31-2). ⁴⁷ An ABI ranging from 0.70 to such as the proximal superficial femoral artery and the popliteal less than 0.90 indicates mild disease; moderate disease correlates artery at Hunter canal. 24

PAD. 25,26 Normal arteries dilate in response to several different evaluating the diagnostic accuracy of the ABI have demonstrated stimuli, such as acetylcholine, serotonin, thrombin, and bradykinin, that it can differentiate between normal and angiographically as well as shear stress induced by increases in blood flow that diseased limbs with a sensitivity of 97% and a specificity of 100% increase nitric oxide production. 27-30 Endothelium from a subject 48 and that the resting ABI is a significant predictive variable for with PAD is impaired; the production and bioavailability of nitric the severity of angiographic disease. 49 oxide in the artery wall are decreased. 31,32 There is increasing evidence for the participation of inflammatory cells as mediators in claudication, defined as pain in one or both legs when walking, atherogenesis and plaque rupture in arterial disease of all vascular beds. 33

There is growing evidence that several nontraditional risk of cardiovascular disease, being involved at all stages of plaque

Several studies have noted increased levels of inflammatory be associated with increased PAD risk. 5 Multiple studies have markers in subjects with PAD. 13,35 Subjects with PAD have shown elevated levels of inflammatory biomarkers in men and increased platelet activity, platelet hyperactivity (as assessed by women with PAD. ^{13,14} In a community-based sample from the platelet aggregation), mean platelet volume, platelet factor 4, Ş -Framingham Offspring Study, 15 the group of inflammatory thromboglobulin, and P-selectin expression receptors on platelets. biomarkers was related to both ankle-brachial index and clinical 21,36,37 Nevertheless, there is a lack of data comparing the association between platelet markers with different pathophysiological

Other pathological considerations for PAD include altered

PAD has two distinct manifestations, claudication and critical 1). 5

Others have noted differences between large-vessel PAD and

ankle-brachial index (ABI). 40-43 In the supine position, the ankle and arm systolic pressures are approximately the same, and on standing, the ankle systolic pressure is somewhat higher than that of the arm. 44 Thus, in the supine position, the measured ankle systolic blood pressure divided by the brachial systolic blood pressure is normally between 1.0 and 1.3. 45 However, if a fixed obstruction of the arterial lumen is present, as is the case with atherosclerosis rhotic disease, a pressure gradient occurs, resulting in a reduced downstream pressure and concomitant The reduction in the ABI. 44

An ABI < 0.9 is considered abnormal and diagnostic of PAD. with an ABI ranging from 0.40 to less than 0.70; and severe Vasodilator capability is severely decreased in subjects with disease is associated with an ABI of no more than 0.40.40 Studies

Approximately one third of patients with PAD have typical

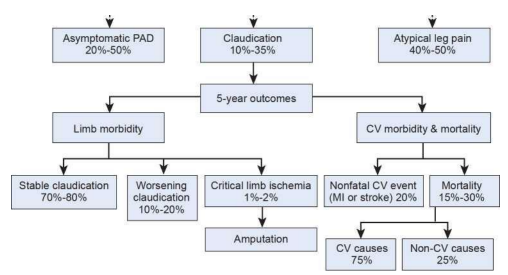


FIGURE 31-1 The natural history of atherosclerotic lower extremity peripheral arterial disease (PAD). Individuals with atherosclerotic lower extremity PAD may be asymptomatic (without identified ischemic leg symptoms, albeit with a functional impairment), present with leg symptoms (classic claudication or atypical leg symptoms), or present with critical limb ischemia. All individuals with PAD face a risk of progressive limb ischemic symptoms as well as a high short-term cardiovascular ischemic event rate and increased mortality. These event rates are most clearly defined for individuals with claudication or critical limb ischemia and less well defined for individuals with asymptomatic PAD. CV, cardiovascular; MI, myocardial infarction. (Modified from Weitz JI, Byrne J, Clagett GP, et al: Diagnosis and treatment of chronic arterial insufficiency of the lower extremities: a critical review. Circulation 94:3026, 1996.)

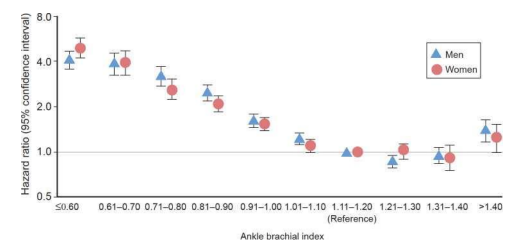


FIGURE 31-2 Hazard ratios for total mortality in men and women by ankle-brachial index in the ABI Collaboration.

primarily affecting the calves, that does not go away with be repeated for the other leg. The lowest ABI between both legs is continued walking and is relieved by rest. 45

Ankle-Brachial Index Technique

the equipment required is inexpensive and portable 40,45,50 (Fig. The pressures in the leg may be supranormal (ABI > 1.4) because of 31-3). The study is done with the patient in the supine position after the inability to compress the artery, especially in patients with resting for at least 5 minutes. The "traditional" method for diabetes. 44 A supranormal pressure does not allow determination conducting the ABI test is to use an ordinary blood pressure cuff of whether an obstructive plaque is present, and thus the and to measure the systolic blood pressure with a Doppler information is considered not diagnostic. If an incompressible ultrasonic velocity signal probe. The blood pressure is measured in artery is found, the patient should be referred to an accredited both arms, and if a discrepancy exists, the higher of the two systolic vascular laboratory for blood pressure values is used. The Doppler probe is then moved over the posterior tibial artery and then over the dorsalis pedis artery to measure the respective ankle pressures. The higher of the two ankle pressures is typically used to calculate the leg ABI. However, recent information indicates that use of the lower of the two may identify more individuals with PAD. 51 The process should

the ABI that stratifies the patient's risk for functional impairment and adverse cardiovascular event. 52,53

Limitations of Ankle-Brachial Index

The ABI can be measured in the office or hospital setting because As with most tests, measurement of the ABI has some limitations.

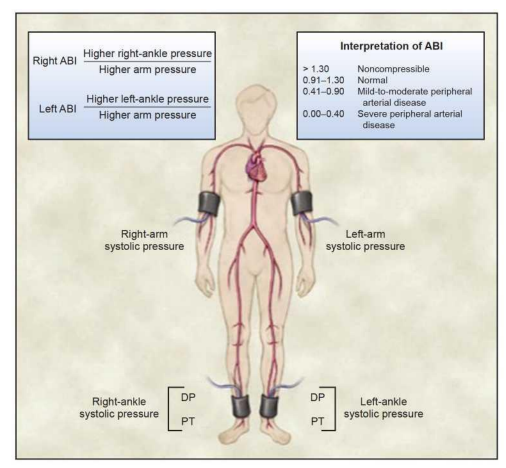


FIGURE 31-3 Ankle-brachial index. DP, dorsalis pedis; PT, posterior tibial artery. (Modified from Hiatt WR; Medical treatment of peripheral arterial disease and claudication. N Engl J Med 344;1608, 2001.)

measurement of a toe-brachial index or other noninvasive testing. Brachial Index

Another problem that may preclude accurate analysis with the ABI is the presence of bilateral subclavian artery stenosis. The result of this hemodynamic occlusion is a false reduction in the "true" systemic circulation systolic pressure and inac clean denominator in the calculation of the ABI. 46 Other limitations of the ABI that should be recognized in considering surgery include its inability to localize arterial lesions accurately 54 and the lack of an association between the ABI and the predicted potential for wound healing. 55

Ankle-Brachial Index in Asymptomatic Peripheral Arterial Disease

Up to two thirds of patients who have a reduced ABI do not have classic symptoms of intermittent claudication. 45,56,57 Individuals with PAD who do not have classic intermittent claudication symptoms have significant functional impairment, functional decline, and cardiovascular events compared with those without PAD. 58 The collaboration of 16 international cohorts including more than 48,000 individuals demonstrated that the ABI provided independent risk information over and above the Framingham risk score, and a low ABI significantly increased the risk of total and cardiovascular mortality and major coronary events across all Framingham risk categories. 47 In fact, the ABI resulted in reclassification of the Framingham risk estimate in approximately 20% of men and one third of women. An analysis of 102 subjects with a recent stroke or transient ischemic attack found that 26% had asymptomatic PAD as detected by ABI measurement. 59 Subjects with asymptomatic PAD had a fivefold greater adjusted increase in cardiovascular events than did subjects without PAD.

Given the wealth of information obtained from this simple hemodynamic test, current American College of Cardiology/ American Heart Association (ACC/AHA) and Inter-Society Consensus for the Management of Peripheral Arterial Disease (TASC II) 38,60 guidelines have provided Class IA recommendation for measurement of the ABI in "at-risk" populations. We recommend the following protocol for measuring the ABI in at-risk populations (Fig. 31-4).

Vascular Exercise Testing

The measurement of the ABI before and after exercise provides additional diagnostic information about the presence and severity of claudication. 40,44,48 The normal response of the ankle systolic pressure is an increase with exercise. However, if significant lower extremity arterial obstruction is present, the ankle systolic pressure may decrease because of a pressure gradient across the blockage while the arm pressure may increase, resulting in reduced ABI.

The ABI will typically return to pre-exercise level within 5 minutes after cessation of exercise. Thus, if clinical suspicion is high that leg discomfort is due to claudication, an exercise test will provide confirmatory information in a patient with "normal" or borderline ABI. Alternatively, if the

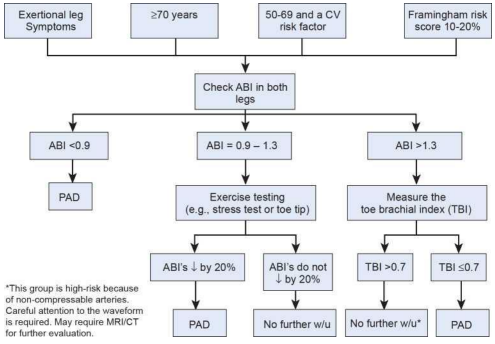


FIGURE 31-4 Flow diagram for measurement of ankle-brachial index.

ABI does not change with exercise, leg discomfort with exercise may be due to another cause, such as spinal stenosis.

The clinical protocol for vascular stress testing may use a standard exercise treadmill or pedal plantar flexion (heel raises). 61,62 One common regimen is for the patient to walk at a standard speed and grade for a predetermined period (ie, 2 mph at 12% incline for 5 minutes) or until claudication develops. 44,46 Immediately after cessation of exercise, the patient is asked to lie in a supine position, and the ankle systolic blood pressure is measured. 44 The pedal plantar flexion exercise test involves the patient's standing facing a wall while using light fingertip support for balance. The patient raises the heels as high as possible while keeping the knees straight and then immediately lowers them; the cycle is repeated 30 to 50 times or until claudication symptoms occur. As with treadmill testing, the ABI is calculated with the patient in a supine position immediately after completing the exercise sequence. 61,62 Postexercise measurement of the ABI may identify an additional 30% of patients with PAD. 63

Segmental Pressures and Pulse Volume Recordings

The ABI, although useful for diagnosis of the presence of PAD, does not provide for the level of disease, an important factor in determining how to treat claudication. A common procedure in many vascular laboratories is measurement of multiple or segmental pressures in the lower extremities along with pulse volume recordings that measure the magnitude and contour of the blood pulse volume. 44,46 This combination of segmental pressures and pulse volume recordings has demonstrated 95% accuracy compared with angiography. 46

An alternative to pulse volume recording measurement is blood velocity waveform analysis, ^{44,46} in which a continuous-wave Doppler probe is used over multiple arterial segments to detect the blood flow velocity and the velocity patterns. ^{44,46} The normal blood flow pattern is triphasic (forward, reverse, and late forward flow), and a change in this pattern to biphasic or monophasic indicates a flow-reducing lesion. ⁴⁴ One major disadvantage of segmental pressure, pulse volume recording, and Doppler velocity waveform analysis is the inability to visualize the anatomy and to pinpoint the artery being studied. ^{44,46} However, the pulse volume recording and Doppler velocity waveform analysis are particularly useful in assessing patients

with supranormal pressures, such as may occur in diabetic patients, due to medial artery calcification. 44,46

Ultrasonic Duplex Scanning

Duplex ultrasound B-mode imaging combined with spectral Doppler analysis is used to localize occlusions more precisely than arterial segments or to more fully characterize the severity and morphology of occlusions. ⁴⁶ The artery characteristics provided from duplex ultrasound imaging include artery wall thickness, degree of flow turbulence, vessel morphology, and changes in blood flow velocity in areas of stenosis. ⁴²

Compared with x-ray contrast angiography, the accuracy (specificity) of duplex ultrasound is very high (92% to 98%), although its sensitivity for assessment of stenosis is variable, depending on the size of the vessel. ⁴²

Some data indicate that the sensitivity for stenosis measurement with duplex ultrasound is lower for smaller arteries than for larger arteries in the limb. The appropriate applications for duplex ultrasound include preparation for planned angioplasty or surgical procedure, detection of restenosis after an endovascular procedure, and surveillance of femoro popliteal or distal saphenous vein grafts for detection of myo-intimal lesions before graft failure. 40,44

If the specific anatomical location and further assessment of stenosis are warranted, then other noninvasive imaging techniques such as spiral computed tomography angiography and magnetic resonance angiography may be used in addition to or instead of duplex ultrasonography. ⁴⁶ Although non-invasive imaging studies are becoming more commonly used preoperatively, catheter-based angiography is still considered the "gold standard."

FORECAST

Symptomatic and asymptomatic PAD is associated with an increased risk for morbidity and mortality. Pooled data from 11 studies in six countries found that PAD, defined by a low

ABI (< 0.9), was associated with an increased risk of subsequent all-cause mortality (RR, 1.60), cardiovascular mortality (RR, 1.96), coronary heart disease (RR, 1.45), and stroke (RR, 1.35) after adjustment for age, sex, conventional cardiovascular risk factors, and prevalent cardiovascular disease. An analysis from the Reduction of Atherothrombosis for Continued Health (REACH) registry 64 spanning 44 countries demonstrated that patients with established atherosclerosis in more than one vascular bed had substantially higher event rates than did patients with atherosclerotic disease in only one vascular bed. In this cohort of approximately 68,000 patients, the annual rate of myocardial infarction, stroke, or death from cardiovascular causes for patients with PAD was 5%. 64

A recent prospective cohort demonstrated a similar high risk of mortality in symptomatic and asymptomatic patients with PAD, and it was significantly higher than in those without PAD. ⁶⁵ Data from the Clopidogrel for High Atherothrombotic Risk and Ischemic Stabilization, Management, and Avoidance (CHARISMA) trial showed an increased risk of all-cause death, cardiovascular death, myocardial infarction, and stroke in subjects with PAD versus no PAD and found no difference between symptomatic and asymptomatic PAD. 66 In a longitudinal study of older men with diabetes without symptomatic PAD, 14-year mortality was significantly higher in subjects with asymptomatic PAD versus no PAD. In the multivariate analysis of the 14-year follow-up, PAD and diabetes were both associated with an increased risk of death, whereas PAD but not diabetes was associated with increased cardiac events and cardiovascular mortality. 67 Data from the ARIC study between 1987 and 2001 found that for every 0.10 decrease in the ABI, the risk for coronary heart disease increased by 25% in white men, by 20% in white women, by 34% in African American men, and by 32% in African American women. 68

In a recent collaboration of 16 international cohorts, ⁴⁷ the ABI provided independent risk information over and above the Framingham risk score. The hazard ratios for all-cause mortality at different levels of ABI compared with a reference ABI of 1.11 to 1.20 in all studies combined formed a reverse J-shaped curve for both men and women (see Fig. 31-2). For levels of ABI < 1.1, the risk of death increased in a step wise manner with decreasing ABI. An ABI > 1.40 was associated with an increased risk of death for men and women. Similar results were noted for cardiovascular mortality and major coronary events. 47

TREATMENT

The treatment of PAD has evolved during the past decade to include a broad approach, focusing on reducing the risk of the major factors associated with the development and progression of PAD. 45,69 Furthermore, because PAD subjects are at high risk for coronary and cardiovascular events and mortality, 70 much emphasis is placed on the reduction of cardiovascular risk. In fact, the National Cholesterol Education Program (NCEP) Expert Panel on Detection, Evaluation, and Treatment of High Blood Cholesterol in Adults (Adult Treatment Panel III) considered PAD a coronary heart disease risk equivalent, thereby elevating it to the highest risk category. 71 Nevertheless, patients with PAD are undertreated with regard to the use of lipid-lowering and antiplatelet drugs compared with patients with coronary artery disease. 72,73

Smoking Cessation

A plethora of evidence exists that smoking is a very significant risk factor for the incidence of PAD and its consequences. 5.74 Cessation of cigarette smoking is associated with a lower amputation rate, a lower incidence of rest ischemia, and an improvement in maximal treadmill walking distance. 75 Among subjects with PAD who are not smokers, there is a lower rate of

myocardial infarction and mortality than among PAD subjects who do smoke. Furthermore, PAD subjects who discontinue smoking have an improved 5-year sur vival versus that of those who continue to smoke. 75 Physician advice must be the cornerstone of this strategy but should be in conjunction with other proven remedies, including group counseling sessions and pharmacological interventions (nicotine replacement therapy, bupropion, and varenicline).

Lipid-Lowering Therapy

Although dietary modification is often the initial treatment for cholesterol concentration, the addition pharmacotherapies may be necessary to achieve target total 31 cholesterol, HDL-C, and LDL-C levels. All patients with PAD of any severity should achieve an LDL-C concentration of less than 100 mg/dL, whereas a target of less than 70 mg/dL is reasonable in subjects with PAD and atherosclerosis in other arterial beds.

In the Scandinavian Simvastatin Survival Study (4S), simvastatin (20 to 40 mg/day) significantly reduced the incidence of intermittent claudication from 3.6% to 2.3% during a median period of 5.4 years in 4444 patients with prior myocardial infarction or angina and a baseline plasma total cholesterol concentration between 212 and 309 mg/dL (relative risk reduction, 0.62%; 95% CI, 0.44%-0.88%). ⁷⁶ In subjects with established PAD, statin therapy may reduce the incidence of cardiovascular events. In a subgroup analysis of 6748 subjects with PAD from the Heart Protection Study, simvastatin 40 mg daily was associated with a reduction in cardiovascular events, regardless of the presenting cholesterol levels. 77 Statin therapy also improves pain-free walking time. 78 Although statin drugs have the strongest data support in use, cholesterol lowering by other means is effective in decreasing cardiovascular events as well. 79

Blood Pressure Control

Antihypertensive therapy is effective at reducing cardiovascular events in subjects with PAD. 45,80 Concern has been raised about the use of beta blockers in the treatment of hypertension among patients with intermittent claudication, but data do not elicit such fears. In fact, a meta-analysis of 11 studies of beta blocker therapy in patients with intermittent claudication found no significant impairment on walking capacity. 81 As a result, these drugs are not contraindicated in patients with PAD. There is some evidence that angiotensin converting enzyme inhibitor therapy may improve cardiovascular events and increase walking distance in selected patients with PAD. In the Heart Outcomes Prevention Evaluation (HOPE) trial, ramipril (5 to 10 mg/day) decreased cardiovascular events in subjects with PAD. 82

Regarding drug choice, all drugs that lower blood pressure are effective at reducing the risk of cardiovascular events. According to the TASC II guidelines, thiazide diuretics and angiotensin-converting enzyme inhibitors should be considered as initial blood pressure-lowering drugs in PAD to reduce the risk of cardiovascular events. 60

Antiplatelet Therapy

The Antithrombotic Trialists' Collaboration, a systematic overview of 135,000 high-risk patients from 287 trials, demonstrated a reduction in myocardial infarction, stroke, and death with antiplatelet therapy in patients at risk for cardio vascular events. 83 Among 42 PAD trials that included 9214 patients and used a variety of antiplatelet agents, compared with placebo, antiplatelet therapy (combining all agents)



ABI < 0.90 N=314	0.81 (0.58-1.14)
ABI 0.91-0.99 N=324	1.28 (0.86-1.91)

0.0 1.0 2.0

Aspirin better Placebo better

Test for heterogeneity -0.089

FIGURE 31-5 The hazard ratio of adverse cardiovascular events (myocardial infarction, stroke, cardiovascular death, or amputation for chronic limb ischemia) in subjects with an ABI of 0.9 and an ABI between 0.91 and 0.99 according to treatment with aspirin 100 mg/day versus placebo in the POPADAD trial.

demonstrated a significant 23% reduction in the odds of cardiovascular events. However, nearly two thirds evaluated nonaspirin antiplatelet agents, questioning whether the overall benefit of antiplatelet therapy in PAD may have been driven by therapeutic regimens other than aspirin.

Subsequently, the prevention of progression of arterial disease and diabetes (POPADAD) trial found no benefit of aspirin 100 mg/day in diabetic subjects with asymptomatic PAD (ABI < 1.0). ⁸⁴ However, a subgroup analysis from the POPADAD trial demonstrated a borderline interaction, suggesting a greater benefit in those with an ABI < 0.9 (Fig. 31-5). A recent meta-analysis of randomized trials of aspirin versus placebo in participants with PAD did not show a significant reduction in cardiovascular events with aspirin. ⁸⁵ However, aspirin was associated with a 35% reduction in the incidence of non-fatal strokes. ⁸⁵

Several hypotheses have emerged as to why aspirin was not found to significantly decrease cardiovascular events in subjects with PAD, including insufficient power, wrong dose, the variation in patient phenotype studied, and perhaps that PAD represents a diffuse form of atherosclerosis with a high inflammatory burden and platelet activity, which may be less responsive to aspirin. The overall benefit of antiplatelet therapy in the Antithrombotic Trialists' Collaboration was driven by data from trials using picotamide, dipyridamole, and ticlopidine. Importantly, aspirin has been shown to have other benefits, including delay in the rate of progression, reduction of the need for intervention, and reduction of graft failure in patients who have undergone revascularization procedures. ^{86,87}

The Clopidogrel versus Aspirin in Patients at Risk of Ischaemic Events (CAPRIE) trial demonstrated that clopidogrel (75 mg/day) had a modest although significant advantage over aspirin (325 mg/day) for the prevention of cardiovascular events in 19,185 patients with a recent stroke, myocardial infarction, or PAD. 88 Overall, there was a 9% relative risk reduction for cardiovascular events, yet among the subset of patients with PAD, clopidogrel resulted in 23% fewer car diovascular events compared with aspirin. However, the CHARISMA trial found no significant benefit of dual anti platelet therapy with aspirin plus clopidogrel versus aspirin alone in patients with established coronary artery disease, cerebrovascular disease, or PAD as well as in patients with multiple atherosclerotic risk factors. 89

Although a benefit of dual antiplatelet therapy versus aspirin alone was noted in subjects with prior myocardial infarction, ischemic stroke, or symptomatic PAD, ⁹⁰ no reduction in cardiovascular events was noted in the subgroup of subjects with PAD. ⁶⁶ In a prospective randomized trial, picotamide significantly reduced mortality in diabetic patients with PAD compared with aspirin. ⁹¹ This may reflect its greater potency, given a dual mechanism of action through inhibition of platelet

thromboxane A 2 synthase and antagonism of thromboxane A 2 receptors. Future studies of this compound are warranted.

Current guidelines ^{5,60} recommend an antiplatelet agent in subjects with PAD, such as aspirin or clopidogrel. On the basis of the limitations of data available, recommendations for aspirin as an important therapeutic tool for secondary prevention in patients with PAD should not be modified. To best inform evidence-based clinical practice guidelines, more high-quality clinical trials are needed.

Anticoagulation Therapy

Oral anticoagulation with warfarin has not been established to reduce cardiovascular events in patients with PAD because it is no more effective than antiplatelet therapy and confers a higher risk of bleeding. ⁹² The Department of Veterans Affairs Cooperative Study tested combined oral anticoagulation and antiplatelet therapy in patients with PAD. ⁹³ No significant difference was found between groups, and there were 133 deaths in the combined treatment group and 95 deaths in the group receiving aspirin alone.

More recently, the Warfarin and Antiplatelet Vascular Evaluation (WAVE) trial randomized PAD patients to combination therapy with an antiplatelet agent and an oral - anticoagulant agent (target international normalized ratio, 2.0 to 3.0) or to antiplatelet therapy alone. ⁹⁴ In this trial, the - combination treatment arm was not more effective than antiplatelet therapy alone in preventing major cardiovascular complications and was associated with a more than threefold increase in life-threatening bleeding. ⁹⁴

Other Pharmacotherapy

In addition to treatment of cardiovascular risk factors and coexisting diseases to prevent cardiovascular events (myocar dial infarction, stroke, and death) associated with atherosclerosis, therapies exist to provide a significant reduction or elimination of PAD symptoms. Claudication drug therapy for relief of symptoms may involve drugs different from those that would be used for risk reduction (an exception may be lipid-lowering therapy). ⁷⁸

Cilostazol is a phosphodiesterase type 3 inhibitor with properties that inhibit platelet aggregation and vascular smooth muscle proliferation and improve the lipid profile and vasodilation. A meta-analysis of six randomized trials demonstrated that cilostazol improved maximum walking distance and pain-free walking distance. 95 An advisory from the US Food and Drug Administration stated that cilostazol should not be used in patients with congestive heart failure because other phosphodiesterase type 3 inhibitors have been demonstrated to worsen survival in this cohort. The effect of cilostazol on cardiovascular morbidity and mortality remains unknown.

Pentoxifylline is a xanthine derivative used to treat patients with intermittent claudication. Its mechanism of action is thought to be a rheological modifier—increase in red blood cell deformity and decreases in fibrinogen concentration , platelet adhesiveness, and whole-blood viscosity. A meta-analysis demonstrated a modestly improved walking distance, substantially less effective than either cilostazol or a supervised exercise program. ⁹⁶

Exercise

For patients with symptomatic PAD, exercise therapy is a key component of reducing symptoms. A supervised program of treadmill-based walking exercise can induce a training



response characterized by large improvements in treadmill exercise performance, peak oxygen consumption, endothelial function, and quality of life. ^{5,60} Exercise is more effective than angioplasty for improvement of walking time and is also more effective than antiplatelet therapy, but it does not differ significantly from surgical treatment. ⁹⁷ Possible mechanisms underlying the exercise response in PAD include improvements in endothelial function, skeletal muscle metabolism, and blood viscosity and a reduction in systemic inflammation . Although it is less well studied, exercise may also improve survival. ⁹⁸ In a prospective observational study of 225 men and women with PAD in whom physical activity was measured with a vertical accelerometer, individuals in the highest quartile of measured activity had a significantly lower mortality than those in the lowest quartile (hazard ratio, 0.29; 95% CI, 0.10-0.83).

Dietary Intervention

The dietary approach for treatment of PAD is the same as for patients with coronary artery and carotid artery atherosclerosis rotic disease. Patients are advised to balance calorie intake and physical activity to achieve and maintain a healthy body weight; to consume a diet rich in vegetables and fruits; to choose wholegrain, high-fiber foods; to consume fish, especially oily fish, at least twice a week; and to limit intake of saturated fat to < 7% of energy, *trans* -fat to < 1% of energy, and cholesterol to < 300 mg/day. ⁹⁹

REVASCULARIZATION FOR PERIPHERAL ARTERIAL DISEASE

The treatment approach with revascularization for PAD involves two distinct populations, those with stable claudication and those with critical limb ischemia. According to the ACC/AHA guidelines, for patients with stable claudication, endovascular or surgical revascularization is "indicated for individuals with a vocational or lifestyle-limiting disability due to claudication when clinical features suggest a reason able likelihood of symptomatic improvement with endovas cular intervention and (a) there has been an inadequate response to exercise or pharmacological therapy and/or (b) there is a very favorable risk-benefit ratio (eg, focal aortoiliac occlusive disease)" (Fig. 31-6). In contrast, those patients with critical limb ischemia deserve consideration for immediate revascularization.

The Inter-Society Consensus for the Management of PAD (TASC) was produced to assess the evidence for PAD diagnosis and treatment and to help guide clinicians in their care of patients with PAD. ⁶⁰ The TASC recommendation regarding determination of the best method of revascularization for treatment of claudication is based on the balance between the risk of a specific intervention and the degree and durability of the improvement that can be expected from the intervention. The outcome of the revascularization procedure depends on the anatomical and clinical features of the lesion.

The anatomical features that have an impact on vessel patency after procedures are the severity of disease in run-off arteries (those distal to the treated lesion), the length of the stenosis or occlusion, and the number of lesions treated. The major clinical factors that affect the outcome of revascularization procedures include diabetes mellitus, smoking, renal disease, and severity of ischemia. The TASC classification of arterial lesions includes A lesions, which should be treated with an endovascular approach; B lesions, which also should be treated with an endovascular approach unless an open revascularization is required for other associated lesions in the same anatomical area; C lesions, for which open revascularization produces superior results and endovascular treatment should 519 be reserved for high-risk patients for surgery; and D lesions, for which the open surgical approach is preferred (Tables 31-1 and 31-2).

A detailed approach to revascularization is provided in the ACC/AHA and TASC II guidelines. 5,60 In brief, the endovas cular treatment of iliac lesions results in the most durable outcome, whereas infrainguinal disease is much more problematic. There are only small clinical studies available directly comparing endovascular treatments with best medical management. The CLEVER (Claudication: Exercise Versus I Endoluminal Revascularization) study, funded by the National Institutes of Health's Heart, Lung, and Blood Institutes, is a prospective, multicenter, randomized, controlled | clinical trial evaluating the relative efficacy, safety, and health 31 economic impact of endovascular and noninvasive treatment strategies for people with aortoiliac PAD and claudication. The results of the CLEVER trial are anticipated to provide needed comparative evidence for revascularization and non-invasive medical approaches. Future studies comparing an invasive versus medical therapy approach are warranted in other anatomical areas, including the femoral and more distal arteries.

ABDOMINAL AORTIC ANEURYSM

An abdominal aortic aneurysm (AAA) is considered to be present when the minimum anteroposterior diameter of the aorta reaches 3.0 cm. ⁵ However, this primary definition does not account for the wide variation in body size, so other definitions have emerged, including a 50% increase in size relative to the proximal normal segment. ^{100,101}

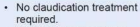
During the last decade, AAA has increasingly been recognized as an important cause of mortality in older persons. In 2006, for example, AAA was noted to be the 13th and 14th leading cause of mortality in the United States among subjects 60 to 69 years and 70 to 79 years, respectively. ¹⁰²

Epidemiology

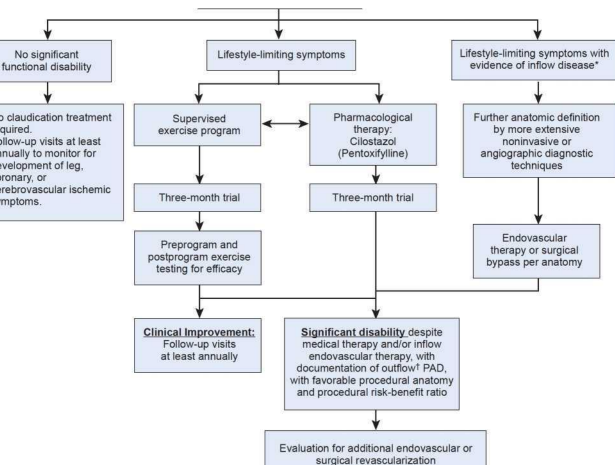
The most important risk factors for AAA are increasing age, smoking, and male sex. ¹⁰³⁻¹⁰⁵ Other factors share many of the same coronary risk factors, including race, atherosclerosis, hypertension, and family history. ^{103,106-111} A recent analysis from the Women's Health Initiative confirmed the importance of smoking and age as robust risk factors in women. ¹¹² As in men, a negative association between diabetes and AAA was reported, suggesting a common pathophysiological mechanism of AAA in men and women. ¹¹³

A series of 46,000 autopsies done in Sweden found an age-standardized AAA prevalence rate of approximately 4.7% among men and 3.0% among women. The prevalence among men increased rapidly after the age of 55 years and reached a peak of 5.9% at the age of 80 years; the prevalence among women increased after the age of 70 years and reached a peak of 4.5% above the age of 90 years. ¹¹⁴ Because AAAs are easily detectable with ultrasonography, more recent studies have evaluated the prevalence of AAAs in the community.

In the final result of the Aneurysm Detection and Management (ADAM) study screening program of the Department of Veterans Affairs, which screened more than 126,196 veterans aged 50 to 79 years, investigators reported a prevalence of 4.2% for aneurysms 3.0 cm or larger and 1.3% for aneurysms 4.0 cm or larger. ¹⁰⁹ In general, the prevalence of AAAs 2.9 to 4.9 cm in diameter ranges from 1.3% for men aged 45 to 54 years up to 12.5% for men 75 to 84 years of age. Comparable prevalence figures for women are 0% and 5.2%, respectively. ⁵



Follow-up visits at least annually to monitor for development of leg. coronary, or cerebrovascular ischemic symptoms.



Confirmed PAD diagnosis

FIGURE 31-6 Treatment of claudication. Inflow disease should be suspected in individuals with gluteal or thigh claudication and femoral pulse diminution or bruit and should be confirmed by noninvasive vascular laboratory diagnostic evidence of aortoiliac stenoses. Outflow disease represents femoropopliteal and infrapopliteal stenoses (the presence of occlusive lesions in the lower extremity arterial tree below the inguinal ligament from the common femoral artery to the pedal vessels). (Reproduced with permission from Hirsch AT, Haskal ZJ, Hertzer NR, et al: ACC/AHA 2005 Practice Guidelines for the management of patients with peripheral arterial disease [lower extremity, renal, mesenteric, and abdominal aortic]: a collaborative report from the American Association for Vascular Surgery/Society for Vascular Surgery, Society for Cardiovascular Angiography and Interventions, Society for Vascular Medicine and Biology, Society of Interventional Radiology, and the ACC/AHA Task Force on Practice Guidelines [Writing Committee to Develop Guidelines for the Management of Patients With Peripheral Arterial Disease]: endorsed by the American Association of Cardiovascular and Pulmonary Rehabilitation; National Heart, Lung, and Blood Institute; Society for Vascular Nursing; Trans-Atlantic Inter-Society Consensus; and Vascular Disease Foundation. J Am Coll Cardiol 47:1239, 2006.)

Abdominal Aortic Aneurysm Screening

The evaluation for an AAA involves physical examination and abdominal ultrasonography. The bifurcation of the aorta occurs at the level of the umbilicus, and palpation may detect a pulsatile mass in the abdomen. Palpation of AAA appears to be safe and has not been reported to precipitate rupture. 115 The physical detection of AAA is significantly hampered by obesity but nevertheless should be included on all vascular examinations. Abdominal palpation will detect most AAAs large enough to warrant surgery, but it cannot be relied on to exclude the diagnosis. 115 The most reliable screening method for AAA is ultrasonography because of high sensitivity (95% to 100%) and specificity (nearly 100%) as well as safety and relatively low cost. 116 Other imaging modalities on which AAA may be seen are abdominal radiography, abdominal computed tomography, and magnetic resonance imaging, but they are not considered first-line evaluation for AAA.

The Multicenter Aneurysm Screening Study (MASS) evaluated a population-based sample of 67,800 men between the ages of 65 and 74 years. 117 subjects were randomly assigned to receive an invitation to undergo ultrasound screening or no correspondence. Of the 33,839 men invited to undergo screening, 1333 aneurysms were detected (prevalence of 4.9%). Ultrasound examination was repeated annually in those with an aortic diameter of 3.0 to 4.4 cm and every 3 months in those with an aortic diameter of 4.5 to 5.4 cm. Those patients with an aortic diameter > 5.5 cm, an increase in a ortic diameter by more than 1 cm in a year, or symptoms attributed to the aneurysm were referred to surgery. Surgical repair was performed significantly more often in the screened group (354 patients) versus the control group (146 patients). Overall, a substantial reduction in aneurysm-related mortality could be

TABLE 31-1	TASC Classification of Aortoiliac Lesions
Type A lesions	Unilateral or bilateral stenoses of CIA
	Unilateral or bilateral single short (< 3 cm) stenosis of EIA
Type B lesions	Short (< 3 cm) stenosis of infrarenal aorta Unilateral CIA occlusion Single or multiple stenosis totaling 3-10 cm involving the EIA not extending into the CFA Unilateral EIA occlusion not involving the origins of internal iliac or CFA
Type C lesions	Bilateral CIA occlusions Bilateral EIA stenoses 3-10 cm long not extending into the CFA Unilateral EIA stenosis extending into the CFA Unilateral EIA occlusion that involves the origins of internal iliac and/or CFA Heavily calcified unilateral EIA occlusion with or without involvement of origins of internal iliac and/or CFA
Type D lesions	Infrarenal aortoiliac occlusion Diffuse disease involving the aorta and both iliac arteries requiring treatment Diffuse multiple stenoses involving the unilateral CIA, EIA, and CFA Unilateral occlusions of both CIA and EIA Bilateral occlusions of EIA Iliac stenoses in patients with AAA requiring treatment and not amenable to endograft placement or other lesions requiring open aortic or iliac surgery

AAA, abdominal aortic aneurysm; CFA, common femoral artery; CIA, common iliac artery; EIA, external iliac artery.

Modified from Norgren L, Hiatt WR, Dormandy JA, et al: Inter-Society Consensus for the Management of Peripheral Arterial Disease (TASC II). *J Vasc Surg* 45(Suppl S):S5, 2007

TABLE 31-2	
	TASC Classification of Femoral Popliteal Lesions
Type A lesions	Single stenosis < 10 cm in length Single occlusion < 5 cm in length
Type B lesions	
	Multiple lesions (stenoses or occlusions), each < 5 cm Single stenosis or occlusion < 15 cm not involving the infrageniculate popliteal artery Single or multiple lesions in the absence of continuous tibial vessels to improve inflow for a distal bypass Heavily calcified occlusion < 5 cm in length
	Single popliteal stenosis
Type C lesions	Multiple stenoses or occlusions totaling > 15 cm with or without heavy calcification Recurrent stenoses or occlusions that need treatment after two endovascular interventions
Type D lesions	Chronic total occlusions of CFA or SFA (> 20 cm, involving the popliteal artery) Chronic total occlusion of popliteal artery and proximal trifurcation vessels

CFA, common femoral artery; SFA, superficial femoral artery.

Modified from Norgren L, Hiatt WR, Dormandy JA, et al: Inter-Society Consensus for the Management of Peripheral Arterial Disease (TASC II). *J Vasc Surg* 45(Suppl S):S5, 2007.

achieved by the implementation of a population screening program. The risk for dying of an AAA during 4.1 years was reduced from 3.3 per 1000 to 1.9 per 1000. 117

For AAA ultrasound screening, the ACC/AHA guidelines recommend the following:

- Men 60 years of age or older who have either siblings or offspring of patients with AAAs should undergo physical examination and ultrasound screening for the detection of aortic aneurysms.
- Men who are 65 to 75 years of age who have ever smoked should undergo a physical examination and one-time ultrasound screening for detection of AAAs.

In January 2007, Medicare coverage for a one-time ultra sound study was made available for Medicare recipients who meet the following criteria:

- Referral from an initial "Welcome to Medicare" physical examination within 6 months of Medicare eligibility.
- Males between 65 and 75 years of age who smoked at least 100 cigarettes.
- Male or female with a family history of AAA.

A study that included 17,540 patients (10,012 women and 7528 men) from 100 hospitals and clinics across the United States reported that AAA occurred in 3.9% of the men and 0.7% of the women. ¹¹⁸ However, certain subgroups of women were at higher risk; women older than 65 years had a fourfold increased odds of having an AAA; women with a history of smoking or a history of heart disease had triple the risk of AAA. These data raise the question of whether certain high-risk groups of women should undergo AAA screening. Only adequately powered outcome studies will help provide therapeutic decisions.

Treatment of Abdominal Aortic Aneurysm

The treatment of AAA involves watchful waiting, open surgical repair, or endovascular stent graft placement. A system attic review of how to manage asymptomatic medium-sized (4.0 to 5.5 cm) aneurysms reported no improvement in survival with early surgical repair. ¹¹⁹ An elective repair is considered for AAA of 5.5 cm in diameter, for those that increase in diameter by more than 0.5 cm within a 6-month interval, or for those that are symptomatic (tenderness or abdominal or back pain). ⁵However, the risk of intervention must be weighed against the potential benefit. The perioperative mortality rate for surgical aortic aneurysm repair ranges from 2.7% to 5.8%. The factors that increase risk include emergent surgery due to rupture, advanced age, chronic kidney disease, cirrhosis, and cardiopulmonary disease. ¹²⁰⁻¹²²

Endovascular repair of AAA is a potential alternative to open surgical repair, but the precise role of endografts in clinical practice has yet to be completely defined. The short term morbidity and mortality of endografts compare favorably with surgical resection. The complications from stent graft placement include endovascular leaks (persistent blood flow into the aneurysmal sac after device placement), device migration, device failure (eg, stent frame fracture), and post implantation syndrome. Because of these potential complications, patients who have undergone endovascular repair of AAA require diligent follow-up with imaging studies annually to evaluate the status of the graft.

A recent randomized, multicenter clinical trial of 881 vet erans with eligible AAA who were candidates for both elective endovascular repair and open repair demonstrated a lower 30-day mortality rate in the endovascular repair group (0.5% versus 3.0%; P = 0.004) without any significant difference in mortality at 2 years (7.0% versus 9.8%; P = 0.13). ¹²³ Longer term outcome data are needed to determine whether endovascular repair is preferable to open surgical repair. ¹²⁴

Prognosis

The rupture of AAA has a mortality rate as high as 90%, 125-127 yet most AAAs never rupture. For that reason, considerable attention has been given to the decision of when and for whom to intervene for elective repair of an AAA and by what



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Study	Туре	Patient Population	Number	Follow-up	Death
Veterans Affairs Cooperative Study(2002)	Cohort	VA setting; AAA >5.5 cm for which repair was not planned	198	1.5 years	57% overall 1 year: 29.8% 2 year: 55.5%
		(contraindication or refusal)			3 year: 75.4%
DREAM (2004)	Randomized controlled trial	Netherlands and Belgium; AAA >5 cm eligible for both endovascular and open repair	Endovascular (n = 171) Open (n = 174)	30 days 2 years	Endovascular vs open Perioperative: 1.2% vs 4.6% (<i>P</i> = 0.10) 2-year: 10.3% vs 10.4% (<i>P</i> = 0.86)
EVAR 1 (2004)	Randomized controlled trial	United Kingdom; men and women >60 years with AAA >5.5 cm eligible for both endovascular and open repair	Endovascular (n = 543) Open (n = 539)	30 days 4 years	Endovascular vs open 30-day: 1.7% vs 4.7% (<i>P</i> <0.05) 4-year: 18.4% vs 20.2% (<i>P</i> = 0.46)
EVAR 2	Randomized controlled trial	United Kingdom; men and women >60 years with AAA >5.5 cm and unfit for open repair	Endovascular (n = 166) No intervention (n = 172)	3.3 years	Endovascular vs open Long term: 44.6 vs 39.5% (P = 0.10)
Medicare (2008)	Cohort	Medicare patients >67 years with a discharge diagnosis of AAA without rupture and a procedural code for open surgical repair or endovascular repair	Endovascular (n = 22,830) Open (n = 22,830)	4 years	Endovascular vs open Perioperative: 1.2% vs 4.8% (P <0.001) (Benefit from endovascular repair persisted for more than 3 years, after which the survival rates associated with the two procedures were similar)
OVER (2009)	Randomized controlled trial	VA setting; eligible for both endovascular and open repair for AAA >5 cm, or associated iliac aneurysm >3 cm, or AAA >4.5 cm with rapid enlargement or saccular morphology	Endovascular (n = 444) Open (n = 437)	1.8 years	Endovascular vs open 30 day: 0.5% vs 3.0% (<i>P</i> = 0.004) 2-year: 7.0% vs 9.8% (<i>P</i> = 0.13)

method. In a surgical cohort of elective and acute intrarenal AAA surgery, mortality within 30 days was approximately fivefold higher for acute AAA surgery, yet no difference was observed during long-term follow-up. 128

The natural history of clinically apparent AAAs of 5.5 cm or more is difficult to determine because most large aneurysms are repaired (Table 31-3). In an observational study from the Veterans Affairs Cooperative Study among 198 patients with AAA of at least 5.5 cm for whom elective repair was not planned because of medical contraindications or patient refusal, investigators reported a 57% mortality rate after a mean followup of 1.5 years. 129 Probable AAA rupture occurred in 23% of the population; the 1-year incidence rates of probable AAA rupture were 9% for AAAs of 5.5 to 5.9 cm, 10% for AAAs of 6.0 to 6.9 cm (19% in the subgroup of 6.5 to 6.9 cm), and 33% for AAAs of 7.0 cm or more. 129

The strongest risk factor for the rupture of an AAA is maximal aortic diameter, and for that reason, it is the dominant indication for repair. A statement from the Joint Council of the American Association for Vascular Surgery and Society for Vascular Surgery estimated the annual rupture risk according to AAA diameter 130:

- 0% for aneurysms < 4.0 cm in diameter
- 0.5% to 5% for those 4.0 to 4.9 cm in diameter
- 3% to 15% for those 5.0 to 5.9 cm in diameter
- 10% to 20% for those 6.0 to 6.9 cm in diameter
- 20% to 40% for those 7.0 to 7.9 cm in diameter
- 30% to 50% for those > 8.0 cm in diameter

Although several biomarkers have been proposed to help identify AAA rupture or expansion, most have weak or no correlation with the clinical course of AAA. 131,132 There are some data that sex may dictate the risk of rupture 133; the rate of rupture of aneurysms that were 4.0 to 5.5 cm in diameter was four times higher in women compared with men. A meta analysis showed that the annual risk of rupture of large AAA (5 cm in diameter) was 18% (95% CI, 8% to 26%) in women versus 12% (95% CI, 5% to 20%) in men. 130,134 Other factors increase the risk of rupture, such as rate of expansion, continued smoking, uncontrolled hypertension, and increased wall stress. 130

Importantly, morbidity and mortality from AAA have improved after elective repair. 135 Analysis of Medicare and Nationwide Inpatient Sample data bases has demonstrated an improvement in 30-day mortality and overall short-term outcomes with endovascular aneurysm repair. 136,137 A recent report from the Swedish aneurysm registry demonstrated an improvement in long-term survival of Swedish citizens treated for AAA with endovascular repair. 138 Data from the United States found similar results-endovascular repair of AAA is associated with a decrease in rupture of AAA and overall patient survival. ¹³⁷Long-term randomized data are essential to properly evaluate current therapy for AAA.

CONCLUSION

PAD, a highly prevalent condition, is most commonly caused by atherosclerotic occlusion of the arteries to the lower extremities, and it is an important manifestation of systemic atherosclerosis. The risk factors for PAD are similar to those for coronary disease, including increasing age, hypertension, dyslipidemia, cigarette smoking, and diabetes mellitus.

increased risk for cardiovascular morbidity and mortality. A simple, inexpensive, noninvasive tool that correlates well with angiographic disease severity and functional symptoms is the ankle-brachial index, and the lower the ankle-brachial index, the greater the risk of 27.

The treatment of PAD has evolved during the past decade to include a broad approach, focusing on reducing the risk of the major 29. Anderson TJ, Uehata A, Gerhard MD, et al.: Close relationship of endothelial function in the human factors associated with the development and progression of atherosclerosis. This risk, along with the severity of claudication in symptomatic patients, can be substantially reduced by targeting the prevalence and level of risk factors by means of lifestyle modification and effective medical therapies. Revascularization for the treatment 32. Sydow K, Hornig B, Arakawa N, et al: Endothelial dysfunction in patients with peripheral arterial of PAD is based on the balance between the risk of a specific intervention and the degree and durability of the improvement that 34. can be expected from the intervention.

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CHAPTER 32

Endothelial Function and Dysfunction

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- KEY POINTS
- Advances in vascular biology and imaging technology have greatly contributed to further elucidation of the complexities of vascular structure and function and the significance of the endothelium in the development of atherosclerosis.
- Vascular endothelium is a vast dynamic paracrine system that regulates several key biologic and molecular functions serving to maintain vascular health and homeostasis.
- The endothelium functions primarily to modulate vascular tone/vasomotion, to maintain an anticoagulant/ profibrinolytic state, to inhibit platelet aggregation and adhesion, to inhibit vascular smooth muscle cell proliferation and migration, and to maintain an antiinflammatory milieu.
- Endothelial activation refers to the biologic response to impairment in vascular homeostasis that engenders a new molecular or functional homeostasis.
- Environmental and genetic factors, such as cardiovascular risk factors, impose an oxidative stress on the vasculature through mechanisms such as the nitric oxide pathway.
- Translational research from experimental discoveries in vascular biology to the clinical arena has spawned applications of new and emerging invasive and noninvasive techniques to measure endothelial function and dysfunction.

 Ongoing research in the development and application of primarily noninvasive imaging techniques to measure endothelial function and dysfunction continues in the pursuit of tests for subclinical disease states, targeting therapeutic strategies and prognosis.

modalities for diagnosis, therapeutics, and

More commonly the arterio-sclerosis results from the bad use of good vessels. William Osler (The Principles and Practice of Medicine, 1892)

Cardiovascular disease incurs a major burden to the public health and health care system. In the past several decades, advances in vascular biology have greatly contributed to our understanding of the complexities of vascular structure and function and the significance of the endothelium in the development of atherosclerosis. Atherosclerosis ubiquitous, complex disease process that is dynamic and multi- factorial. Genetic and various environmental factors, with their complex interactions, lead to the initiation and progression of atherosclerosis during decades. The vascular endothelium, the largest paracrine organ, vastly forms the inner lining of all blood vessels in the vasculature to maintain vascular homeostasis myriad complex biological properties and physiologic processes. Strategically located between the vessel lumen and smooth muscle layer, the endothelium serves to modulate vascular tone, cell growth, plate let and leukocyte interactions, inflamma tion, and thrombogenicity.

Physical and chemical stimuli in the vascular milieu initiate and propagate various physiological and molecular processes through signal transduction mechanisms not yet fully understood and through the elaboration and secretion of various substances. The magnitude and duration of exposure to cardiovascular risk factors impose injury on the vasculature, primarily through oxidative stress mechanisms that promote procoagulation, inflammation, vasoconstriction, and proliferation of cell growth. The complexity of atherosclerosis translates into a continuum of cardiovascular disease processes that manifest derangements in endothelial functions, which begin early in the pathogenesis of disease. The study of endothelium from cell to organ system poses the challenge for development and application of various genetic and chemical biomarkers and imaging

prognosis.

Various invasive and noninvasive imaging modalities have evolved in the last several decades to clinically evaluate the structure and function of various vascular beds in health and disease states. Advances in experimental and clinical research under score the complexity of vascular homeostasis . Perturbations in homeostasis induced environmental factors or the local biological milieu are manifestations of endothelial dysfunction that require further definition of the specific vascular property involved. The term endothelial activation refers to the biological response to impairment in vascular homeostasis that engenders a new molecular functional homeostasis. 1

Imaging techniques to assess endothelial function, particularly vascular vasomotion, emerged concurrently with the advances in vascular biology, coupled with a keen interest in the use of biomarkers and imaging modalities and techniques to detect subclinical disease while targeting preventive and therapeutic interventions. Investigation continues in the clinical applications to enhance early detection and to guide therapy and prognosis.

ENDOTHELIUM AND ENDOTHELIAL PHYSIOLOGY: GENERAL OVERVIEW

Malpighi's discovery in the 17th century of the endothelium as a physical separation between blood and tissue with no substantial functionality persisted through the 19th and mid-20th centuries. The endothelium is a 0.2-to 4-|rm-thick monolayer of squamous endothelial cells lining the entire surface of the vasculature, including endocardium, arteries, arterioles, capillaries, venules,

Vascular tone
Vascular permeability
Vascular remodeling
Anticoagulant, profibrinolytic, and procoagulant activities
Inflammatory and immunopathological responses
Interactions with blood components

Healthy endothelium

Vasodilator: endothelium dependent

Antihypertrophic: inhibits vascular smooth muscle cell proliferation and migration

Anticoagulant and profibrinolytic

Antithrombotic: inhibits platelet adhesion and aggregation
Anti-inflammatory: inhibits leukocyte adhesion and migration

veins, adventitial vasa vasorum, and other microcirculation. ¹ Once thought to be a passive, semipermeable membrane between blood flow and vascular wall, the endothelium is essentially a vast autocrine, paracrine, and endocrine organ that spans a surface area of approximately 700 m ².

The endothelium is strategically located at the interface of the blood circulation, blood components, and vascular smooth muscle and adventitia. As such, the endothelium is the major regulator of vascular homeostasis, which occurs through myriad diverse and interrelated physiological functions. These include the regulation of vasomotion, smooth muscle cell proliferation, inflammation, thrombolysis, homeostasis, platelet aggregation, immune responses, cell proliferation, and free radical production. Normal healthy endothelium exerts a variety of effects to maintain vascular homeostasis of these various biological functions (Table 32-1). It performs these functions through an elaborate array of secretable substances and signal transduction mechanisms in response to a number of different physiological, chemical, and mechanical stimuli within and around its surrounding milieu and external environmental factors. 1-3

One of the pivotal roles of the endothelium is the modulation of vascular tone, caliber, and blood flow in response to neural, humoral, and mechanical stimuli by synthesis and release of various vasoactive substances. ²⁻⁴ Specifically, the endothelium plays a key role in the regulation of hemostasis and thrombosis, vascular tone, inflammation, and vascular growth and remodeling under normal conditions. Laminar stress is among the most important stimuli that help maintain the normal physiological state of the endothelium. This is achieved by reduction of biological activity of proteins by *S*- nitrosylation of cysteine residues and oxidative phosphorylation in mitochondria. ^{5,6} However, during exposure to risk factors such as hypertension, diabetes mellitus, and tobacco smoking, dysfunction of the endothelium ensues, thus disturbing the fine balance and subsequently resulting in atherosclerosis.

Vasomotion

Regulation of vascular tone by endothelium is achieved by generation and secretion of vasoactive substances. The endothelium exerts a significant effect on both vasodilation and vasoconstriction. Endothelium-mediated vasodilation is predominantly achieved by nitric oxide (NO) and prostacyclins (PGI 2). Endothelium-mediated vasoconstriction is regulated by secretion of angiotensin II, platelet-derived growth factors, platelet-activating factor, and endothelin 1, all of which have vasoconstrictive effects. ⁷NO serves as an important vasodilator and forms a major basis for endothelial function and dysfunction.

Prostacyclin, another vasodilator synthesized in the endothelium, is a product of arachidonic acid, which is released from membrane phospholipids in response to shear stress. ^{1,8} Vasodilator effects of prostacyclins are dependent on expression of receptors in vascular smooth muscles. This limits the role of

endothelium-mediated vasodilation in vascular beds where receptors are not expressed. Prostacyclins do not contribute to the maintenance of basal vascular tone of large conduit arteries. Prostacyclins mediate their effect through receptors coupled to adenylate cyclase and elevation of cyclic adenosine monophosphate (cAMP) levels in vascular smooth muscles, of ATP-sensitive potassium channels. stimulation hyperpolarization of cell membrane, and thus inhibition of the development of contraction. Inhibition of the contractile mechanism is also mediated by the expulsion of calcium from the cytosol of vascular smooth muscles. In addition, prosta cyclins contribute to the release of NO by endothelial cells and have a synergistic effect with NO on antiplatelet activity.

Endothelin 1

Endothelin 1 (ET-1) is a potent vasoconstrictor generated in the endothelial cells along with other cell lines. It belongs to a family of structurally related peptides including ET-1, ET-2, and ET-3. 9,10 Mature human ET-1 is derived from a precursor, preproendothelin 1, through an intermediate molecule and requires a peptidase named ET-converting enzyme. This peptidase serves as a critical physiological regulator of ET-1 activity . Shear stress and cyclic stress, along with other factors such as hypoxia, stimulate the generation of ET-1 from endothelial cells. The vasoconstrictor effect of ET-1 is mediated by binding to ET $_{\rm A}$ on vascular smooth muscle cells. The effect on NO and PGI $_{\rm 2}$ generation is mediated by binding to ET $_{\rm B}$ receptors on the endothelium.

Vascular beds exhibit heterogeneity in vasoconstrictor response to ET-1. Whereas the renal endothelium and the coronary endothelium are extremely sensitive to ET-1, the pulmonary circulation exhibits a less sensitive response. In addition to vasoconstriction, ET-1 exhibits a wide range of activities. It has a positive inotropic effect on cardiomyocytes and stimulates the release of atrial natriuretic peptides from atrial myocytes. In addition, it aids in the release of aldosterone and catecholamines from the adrenal cortex and medulla. ET-1 augments the vascular actions of other vasoactive peptides (such as angiotensin II, norepinephrine, and serotonin), participates in leukocyte and platelet activation, and thus facilitates a prothrombotic state. 11,12 It also has an inhibitory effect on renin release from juxtaglomerular cells. Finally, it enhances the release of endothelium-derived relaxing factor and PGI 2 and modulates vascular remodeling. 13

Inflammation

The endothelium, as a result of its location, serves as a potent anti-inflammatory tissue. It is constantly exposed to various pathogens, anti-inflammatory cells, and immunoreactive substances. Disease states or exposures such as hypertension and atherosclerosis constitute inflammatory changes in the vascular wall. Inflammatory changes include altered expression of adhesion molecules. Under shear stress, endothelial cells exhibit a number of anti-inflammatory properties. This response includes prevention of adhesion of circulating inflammatory cells to endothelial cells and release of immunoreactive substances. NO released from endothelial cells aids in limiting leukocyte adhesion. Furthermore, NO has an inhibitory effect on the release of prothrombotic substances, such as von Willebrand factor and P-selection.

NO limits activation of nuclear factor- KB (NF- KB) and thus inhibits adhesion molecule expression and subsequent attachment of immune cells to endothelium. Endothelial



528 dysfunction, as a result of oxidative stress in the endothelial cells, results in activation of NF- K B, a redox-sensitive transcription factor. Subsequent steps include release of chemoattractant proteins such as monocyte chemotactic protein 1 and expression of adhesion molecules (Eselectin, P-selectin, intercellular adhesion molecule 1 [ICAM-1], and vascular cell adhesion molecule 1 [VCAM-1]). This results in monocyte attachment and rolling through the selectins, activation through selectins and chemokines (CCL2, CXCL8, and platelet-activating factor), arrest and adherence through the ■in immunoglobulin G family (ICAM-1, VCAM-1) and integrins (a V P 3), and finally extravasation through platelet/endothelial I cell adhesion molecule 1. Other molecules involved in extravasation of leukocytes include cadherins. ^{14,15}

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Hemostasis and Thrombosis

The endothelium serves as a lining of a compartment that maintains blood flow while allowing the delivery of important nutrients to other organs in the body. It serves this purpose by inhibiting platelet aggregation and clotting cascade activation. It the antithrombotic properties by generating anticoagulants (antithrombin III, thrombomodulin, tissue factor pathway inhibitor, protein C, and heparan sulfate proteoglycans), fibrinolytics (tissue-type plasmino gene activators and urokinase-type plasminogen activators), and platelet inhibitors (NO, prostacyclins, and ADPase [CD39]). On the opposite end of the spectrum, the endothelium maintains hemostasis by production and release of procoagulants (thrombin receptor, protein C receptor, tissue factor, and coagulation factor binding sites) and antifibrinolytics (plasminogen activator inhibitor 1) and by promotion of platelet activation (von Willebrand factor and platelet activating factor).

Whereas normal endothelial cells do not express - procoagulants such as tissue factor, activated endothelial cells rapidly express the same on their cell surface. Similarly, von Willebrand factor is stored in endothelial cells in granules called Weibel-Palade bodies, which are exposed on the endothelial surface in response to injury and other soluble media tors, resulting in formation of a hemostatic plug and platelet adhesion. ¹

Vascular Growth and Remodeling

With better understanding of developmental biology, it is now evident that a close relationship exists between endothelial cells and the hematopoietic cell lineage. It is now known that hemangioblasts may be the common precursor for both cell lineages. The hemangioblasts differentiate into either blood cell precursors or angioblasts, which have endothelial cell precursors. ¹⁶ Vessel growth or angiogenesis after the initial process of development of blood vessels involves the formation of capillary sprouts by endothelial cells. ¹⁷ This process subsequently leads to remodeling and formation of a mature vessel.

Endothelial cells play an important role in the remodeling process during development by secretion of substances that recruit undifferentiated mesenchymal cells and cause their maturation into pericytes or smooth cells. ¹⁸ During endothelial dysfunction, in adult cells, vascular smooth muscle proliferation takes place in the presence of oxidative stress and as a result of decrease in endothelium-derived NO, which has an inhibitory effect on vascular smooth muscle growth.

It is now known that the endothelium along with the vasa vasorum contributes to angiogenesis and neovessel formation in atheromas. These neovessels are instrumental in atherosclerotic plaque progression, instability, and rupture. The microvascular channels formed by invagination of endothelial lumen may also serve for the transport of inflammatory cells, such as leukocytes. This is supported by selectively increased VCAM expression on microvascular endothelial cells. ¹⁹ In addition to leukocyte migration, microvessels cause intraplaque neovascularization, thus leading to hyperpermeability , and result in microhemorrhage and thrombosis. ^{19,20} Evidence of hemosiderin

deposits in the neovascular plexus and their colocalization with thrombotic factors such as von Willebrand factor suggest hemorrhage and thrombosis within the atheroma. Hyperpermeable neovessels allow extravasation of red blood cells. ²¹ Lysis of red blood cells contributes to plaque progression by lipid expansion as their membranes are rich in cholesterol and generation of reactive oxygen species and macrophage activation. ²⁰ As discussed elsewhere, reactive oxygen species deplete endothelium-derived NO and thus limit its inhibitory effect on vascular smooth muscle growth.

Nitric Oxide Pathway

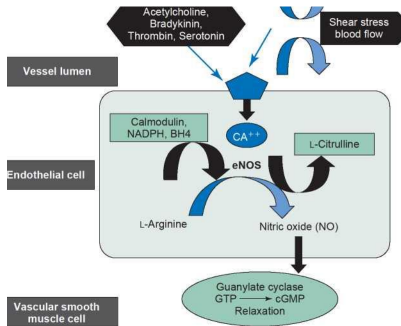
In the early 1980s, Furchgott and Zawadski ²² first demonstrated the obligatory role of the endothelium for vascular relaxation in response to vasoactive substances such as ace tylcholine and postulated the existence of an endothelium-derived relaxing factor, later discovered to be NO. ^{1,23,24} Of all substances secreted or controlled by the endothelium, NO is the most recognized molecule; it is pivotal in maintaining vascular tone and mediates inhibition of coagulation, platelet activation, smooth muscle cell proliferation, and inflammation . Inadequate or lack of NO is implicated in endothelial dysfunction.

NO is produced by the endothelial cells from arginine (Fig. 32-1). It is a heterodiatomic lipophilic free radical that is synthesized by endothelial nitric oxide synthase, a heme containing NAD(P)H-dependent oxygenase that requires cofactors tetrahydrobiopterin and nicotinamide adenine dinucleotide phosphate. ²⁵ Three distinct isoforms of nitric oxide synthase (NOS) are known to exist: endothelial NOS (eNOS), inducible NOS (iNOS), and neuronal NOS (nNOS). All three isoforms belong to a family of arginine hydroxylases. ²⁶ Most of the NO production takes place in the invagina tions of the cell membrane (caveolae) of endothelial cells. It is made from the amino acid -- arginine, which is converted to -- citrulline by means of eNOS. Caveolin 1 regulates the activity of eNOS by binding to calmodulin, with the resulting inhibition of eNOS. However, eNOS activation and NO production are achieved by binding of calcium to calmodulin, causing a displacement of caveolin. ^{27,28}

Activity and synthesis of eNOS are modulated by physiological inhibitors, such as asymmetric dimethylarginine and geranylgeranyl pyrophosphate (intermediate of cholesterol biosynthesis), or pharmacological inhibitors, such as N -monomethyl-1-arginine (1-NMMA) and 1-nitroarginine methyl ester. NO, a potent endogenous vasodilator, is basally secreted in response to various physiological agonists, physical stimuli, and pharmacological agents. It exerts its effect on vascular tone in multiple ways. The basal secretion of NO maintains the vessel in a vasodilatory state and assists in maintaining vascular health through its various antiatherogenic properties.

Once it is secreted, NO diffuses into the subendothelium and exerts a vasodilator effect on vascular smooth muscle cells. It activates the soluble guanylate cyclase, followed by an increase of the intracellular concentration of cyclic guanosine monophosphate. This results in a reduction in the intracellular calcium concentration, followed by a relaxation of vascular smooth muscle cells. ²⁹ In addition, it decreases the expression and activity of the potent vasoconstrictor ET-1. NO has an inhibitory effect on vascular smooth muscle proliferation and migration and extracellular matrix production. ²⁹ NO exerts this effect by inhibiting the activation of NF- K B. This particular attribute contributes to

FIGURE 32-1 Production and release of nitric oxide leading to vasodilation. BH4, tetrahydrobiopterin; Ca ² + , calcium ion; cGMP, cyclic guanosine monophosphate; eNOS, endothelial nitric oxide synthase; GC, guanylate cyclase; GTP, guanosine triphosphate; NADPH, reduced nicotinamide adenine dinucleotide phosphate. (Modified from Behrendt D, Ganz P: Endothelial function. From vascular biology to clinical applications. Am J Cardiol 90:40, 2002.)



anti-inflammatory as well as to antiproliferative properties . ^{30,31} NO exerts its inhibitory effect on platelet activation and aggregation by stimulation of the cAMP pathway. Its effect on the fibrinolysis system is mediated by stimulation of tissue plasminogen activator release. ³² However, the effect on mobilization of progenitor cells and stem cell modulation of their survival and function are areas of immense interest. ³³

Diminished NO activity may be due to four distinct causes: decreased expression of eNOS enzyme, eNOS uncoupling, increased scavenging of NO, and impaired transmission of NO-mediated signaling. ^{34,35} Both physical and humoral stimuli can activate the transcription of eNOS gene. Activation of eNOS by physical stimuli such as shear stress is mediated by Raf, Ras, and ERK1/2 and NF- K B binding to shear stress response element. ³⁶ Humoral factors influencing eNOS transcription include growth factors (vascular endothelial growth factor, basic fibroblast growth factor, epidermal growth factor, and transforming growth factor- §), cytokines, and oxygen radicals.

Uncoupling of eNOS refers to the process by which eNOS switches its predominant function of NO generation to reactive oxygen species formation. When this switch occurs as a result of tetrahydrobiopterin deficiency, generation of oxidants such as superoxide is dominant. However, deficiency of r-arginine also leads to production of hydrogen peroxide. ¹¹ Superoxide anions rapidly interact with NO to form peroxynitrite. Peroxynitrite in turn decreases tetrahydrobiopterin and results in increased consumption and reduced production of NO. ^{35,36} Figure 32-2 summarizes the mechanism of decreased NO bioavailability mediated by reactive oxygen species. In addition, various factors known to regulate eNOS gene expression mediate endothelial function and dysfunction (Table 32-2).

ENDOTHELIAL DYSFUNCTION

No single definition of endothelial dysfunction exists, given the complex and ubiquitous nature of endothelial biology.

Endothelial dysfunction was first described as structural changes or loss of anatomical integrity in the context of atherosclerosis. ³⁷ Endothelial dysfunction includes broad regulatory changes leading to abnormal vasomotion and the expression of a prothrombotic and proinflammatory phenotype of the vascular endothelium. This poses a challenge to define endothelial dysfunction in terms that include all the possible alterations in vascular homeostasis and at various time points in various

disease states.

Endothelial dysfunction encompasses any alterations in the various vascular biology and function, particularly vaso motor function as well as the prothrombotic, proinflammatory, proatherogenic properties. Endothelial activation refers to the biological response to alterations or impairment in vascular homeostasis that would elicit new molecular or functional homeostasis, including gene activation ¹ and the repair response to damaged endothelium. ^{38,39}

Chronic exposure to injury stimuli and reactive oxygen species may surpass the inherent capacity of the endothelium to mount a suitable defense. This may subsequently lead to aging and senescence of the endothelial cells. Finally, the senescent endothelial cells may detach and be released into the circulation.

Activated endothelial cells, apoptotic cells, and their components in the circulation thus may serve as markers for endothelial dysfunction. ⁴⁰The levels of these cell types and their components have been shown to be increased in several inflammatory states, such as coronary artery disease, rheumatoid arthritis, and systemic lupus erythematosus. ^{41,42} Endothelial repair may be due to repopulation of the denuded endothelium by replication of adjacent mature endothelial cells or from circulating endothelial progenitor cells. ⁴³

Endothelial progenitor cells are circulating precursors recruited from the bone marrow, partly by NO-dependent mechanisms. These cells may transform into mature endothelial cells and contribute to the maintenance of endothelial integrity and health. The extent to which this mechanism may be successful depends on the exposure to cardiovascular risk factors and inflammatory stimuli. Under certain circumstances, these cells may transform into macrophages and thus

Hypercholesterolemia, Hypertension, Diabetes, Smoking NAD(P)H oxidase Xanthine oxidase Mitochondria? Reactive oxygen species production 1) BH4 oxidation Inactivation of NO' 0 ADMA f eNOS "uncoupling ONOO eNOS activity 4 NO'4 O 2-f NO O 2' production production

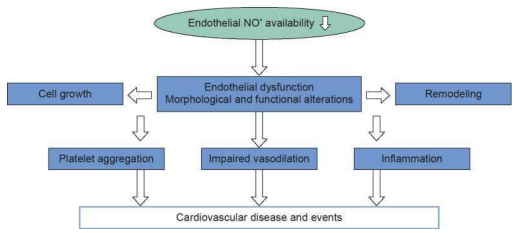


FIGURE 32-2 Schematic representation of mechanisms of decreased NO bioavailability resulting in endothelial dysfunction. Reactive oxygen species decrease the bioavailability of NO in endothelium by three different mechanisms; (1) superoxide reacts with NO to form peroxynitrite anions, causing a decreased bioavailability of NO; (2) reactive oxygen species cause an increased concentration of ADMA, an endogenous inhibitor of eNOS; and (3) uncoupling of eNOS enzyme due to degradation of BH 4. ADMA, asymmetric dimethylarginine; BH 4, tetrahydrobiopterin; DDAH, dimethylarginine dimethylaminohydrolase; eNOS, endothelial NO synthase; NO, nitric oxide. (From Landmesser U, Hornig B, Drexler H: Endothelial function: a critical determinant in atherosclerosis? Circulation 109[Suppl 1]:Il27, 2004.)

may contribute to endothelial dysfunction. Modification of risk factors by interventions such as exercise and statins, as discussed later, may help in mobilization of endothelial progenitor cells and aid in repair. 44,45 Any imbalance between injury and repair may result in endothelial dysfunction. Whereas levels of endothelial progenitor cells may be increased upon exposure to acute insult, chronic inflammatory states result in reduced levels. Reduced levels of endothelial progenitor cells may be a result of overconsumption, inhibited mobilization, or accelerated senescence. Risk factors, by modulating the levels of oxidative stress or NO activity, may influence the levels of circulating progenitor cells. The precise underlying mechanism needs to be understood better.

Oxidative Stress and Endothelial Dysfunction

Oxidative stress refers to a state whereby the rate of formation of reactive oxygen species exceeds the capacity of physiological antioxidant defense mechanisms. Chronic exposure to reactive oxidative species may overwhelm the inherent antioxidant mechanism and thus contribute to endothelial dysfunction or prolonged endothelial activation. Several enzyme systems are sources of free radical production (reactive oxygen species), including NAD(P)H oxidases, nitric oxide synthases, lipoxygenases, cyclooxygenases, oxidoreductases, and mitochondrial oxidases. 46 Balance between production of reactive oxygen species and activity of superoxide dismutase during oxidative phosphorylation is maintained in normal physiological conditions. 47

An abundance of reactive oxidant species alters several important physiological functions, including regulation of blood flow, vasodilation, coagulation, inflammation, and cellular growth, and activates multiple signaling pathways in vascular wall cells that also contribute to the impairments of vascular function and repair. On exposure to risk factors such as obesity and diabetes mellitus, there is an alteration in this balance due to increased substrate delivery, such as circulating free fatty acids. ^{13,45} In coronary artery disease, increased oxidative stress is a result of increased nicotinamide adenine dinucleotide phosphate oxidases and xanthine oxidase. 34,35,46,47 As depicted in Figure 32-2, various risk factors result in increased oxidative stress and contribute to decreased NO bioavailability by three distinct mechanisms. Production of superoxide is increased in the diseased vessels of patients with coronary vascular disease. 48 Superoxide interacts with NO, forming peroxynitrite anions, resulting in consumption of NO and loss of its activity. 48,49

TABLE 32—2 Factors Regulating Endothelial Nitric Oxide Synthase Gene Expression				
Factors	Effect			
Physical factors				
Shear stress	Stimulation			
Hydrostatic pressure	Stimulation			
Humoral factors				
Reactive oxygen species				
Nitric oxide	Inhibition			
Oxidized low-density lipoprotein	Inhibition			
H2O2	Stimulation			
Inflammatory factors				
Tumor necrosis factor- a	Inhibition			
Growth factors				
Vascular endothelial growth factor	Stimulation			
Basic fibroblast growth factor	Stimulation			
Epidermal growth factor	Stimulation			
Transforming growth factor- p	Stimulation			
Peptide hormones Angiotensin II Endothelin 1	Stimulation Stimulation			

Stimulation or inhibition

NO is a key mediator in vascular homeostasis as it serves as an antiatherogenic molecule, promotes vasodilation, and counteracts inflammation, platelet aggregation, and vascular smooth muscle proliferation. A large body of evidence supports the tenet that endothelial dysfunction is caused to some degree by accelerated inactivation of NO by reactive oxygen species, although exact mechanisms are not entirely elucidated and strategies for targeted therapy are still under investigation. Another mechanism is the redox-dependent inhibition of the enzyme dimethylarginine dimethylamino hydrolase. This results in increased concentration of the endogenous eNOS inhibitor asymmetric dimethylarginine (ADMA). Levels of ADMA are elevated in a large number of conditions, including several associated with risk of endothelial dysfunction. 50,51 ADMA concentration may be increased because of impaired excretion or increased synthesis, as in the case of vascular shear stress. 50,51 Accumulation of ADMA acts as a competitive antagonist to the eNOS substrate L-arginine.

Erythropoietin

Role of Endothelial Dysfunction in the Presence of Specific Risk Factors and Atherosclerotic Disease

Several risk factors may play a critical role in contributing to endothelial dysfunction and thus atherosclerosis. Traditional and novel cardiovascular risk factors, including smoking, aging, hyperlipidemia, hypertension, diabetes, and family history of premature atherosclerosis, among others, are associated with a loss or attenuation of endothelium-dependent vasodilation in both children and adults. ^{52,53} Elevated C-reactive protein, chronic systemic infection, obesity, and several immunemediated diseases are also associated with impaired endothelial function. ^{54,55} Given its complex biology, the term *endothelial dysfunction* applies broadly to the various disturbances that contribute over time to the development and clinical expression of atherosclerosis.

The underlying mechanism through which risk factors impart vascular injury initiating endothelial dysfunction is indeed multifactorial, yet it is generally accepted that the predominant mechanism is oxidative stress and redox injury resulting in decreased synthesis or increased degradation of NO. Given the diverse vasculoprotective properties of NO and related pathways, there is increasing evidence to support the notion that endothelial dysfunction begins with impairment of oxidative stress, with subsequent unavailability of NO. Factors that help stimulate the production and release of NO have also been an area of active research.

An important mechanical stimulus that evokes vasodilator release of vasoactive substances, particularly NO, from the vascular endothelium is pulsatile flow and shear stress induced from movement of blood along the endothelial cells. 2-4 Varying degrees of shear stimulus have been shown to have an impact on the degree of vasodilation, which points to the complexity of the underlying vasomotor function, the impact of vascular risk factors and disease states, and the ability to decipher specific 32 mechanisms underlying impaired endothelial vasomotor function. Reactive hyperemia is a transient increase in blood flow, physiologically or mechanically invoked, that immediately follows a designated period of vessel occlusion or flow disturbance. High shear stress on the endothelium provides the stimulus for NO release. In addition, a variety of vasodilators, such as adenosine and hydrogen ions, among others, are released and act locally on the microvessel milieu. 56,5

ENDOTHELIAL VASOMOTOR FUNCTION TESTING

The initial observations and discoveries in vascular biology since the 1980s stimulated intense basic research in vascular biology in the ensuing decades to further delineate the various components and mechanisms of vascular diseases. A vast body of work emerged emphasizing the importance of endothelium-derived NO as a potent endogenous vasodilator that contributes to vascular health in optimal conditions. Moreover, these observations formed the basis for translational research to study the characteristics and clinical significance of endothelium-dependent properties, such as vasodilation, in various vascular beds and in various vascular disease states.

Experimental and clinical techniques using physiological agents and various imaging techniques were developed and applied to study vascular structure and functions of both conduit and resistance vessels in the normal state and within the context of cardiovascular risk factors, such as dyslipidemia , hypertension, diabetes, and early atherosclerosis. Such techniques also spurred further investigations of the impact of therapeutic and lifestyle modifications on endothelial dysfunction and its prognosis. ⁵⁷

Invasive Measures of Coronary Vasoactivity

With the discoveries in endothelial biology in the 1980s, there was growing interest in the potential to clinically examine disturbances in vascular function before the advent of detectable atherosclerosis and clinical events. Numerous studies and methods using various physiological agents and techniques to study endothelium-dependent vasomotor function in conduit and resistance vessels in the normal state and in the presence of cardiovascular risk factors and atherosclerosis ensued. Ludmer and colleagues, ⁵⁸ in the 1980s, first described impaired endothelium-dependent vasomotor function with intracoronary injection of acetylcholine and quantitative coronary angiography in humans with various degrees of atherosclerosis noted by angiography. They demonstrated that like isolated vascular rings used in basic experimentation , human coronaries that appeared angiographically

532 normal or with mild stenosis paradoxically vasoconstricted to acetylcholine but not to nitroglycerin. 58 Subsequently, endothelium-dependent, NO-mediated vasomotor function testing gained notoriety as a useful tool to assess the functional

integrity of vascular endothelium in vivo.

serve as a surrogate marker for the bioavailability of NO. Endothelial function is most commonly measured as the vasomotor response to pharmacological stimuli, such as acetyl choline, methacholine, bradykinin, serotonin, papaverine, and (increased blood flow velocity), exercise, cold pressor test, and mental stress. During cardiac catheterization, alterations in arterial lumen diameter in response to such stimuli 32 can be measured by computerized edge detection software to provide reproducible measurements of the angiographic lumen diameter compared with baseline diameters. Lumen diameter and coronary flow by quantitative coronary angiography and intracoronary Doppler study, respectively, can be used to assess changes in vessel diameter and coronary flow reserve.

Endothelial function in resistance vessels (microcirculation) is also critical in the assessment of endothelial vasomotor function. Resistance vessels regulate blood flow in response to changes in perfusion pressure (autoregulation) and metabolic needs (metabolic regulation). The vascular tone of resistance vessels determines blood flow; therefore, altered blood flow detected by techniques without considerable changes in mean blood pressure indicates changes in vessel tone resistance. Endothelial function of forearm resistance vessels can be adenosine or by forearm blood flow responses to intra-arterial agonists with strain-gauge plethysmography. 59

Angiography is invasive and carries some risk, and it is not conducive to repeated studies in the same individual over time or for the study of relatively low risk populations. Insights from these invasive studies of endothelial vasomotor function and flow reserve provoked a keen interest in the pursuit of similar studies in the peripheral circulation with both invasive and noninvasive techniques. This was motivated by the notion that endothelial dysfunction is both a local and systemic physiological state, along with an interest in detecting the impact of cardiovascular risk factors and the presence of preclinical atherosclerotic disease.

Invasive Assessment of Forearm Microcirculation (Venous Occlusion Plethysmography)

The limitations inherent in coronary artery circulation studies of vasomotion led researchers to pursue the peripheral Technique circulation for further investigation. There was also keen The technique and examination protocol for brachial artery broadly assess the systemic nature of atherosclerosis and the broader impact of cardiovascular risk factors and the potential role to evaluate the impact of interventions and risk factor modifications. Moreover, further study led to an interest in the health and disease.

Venous occlusion plethysmography is an invasive forearm blood flow in response to an intra-arterial infusion of MHz enable visualization a vasoactive substance such as acetylcholine, substance P, or adenosine into either the brachial artery or radial artery or to reactive hyperemia (increased shear stress). The standard testing technique is well described, reliable, and highly reproducible and typically used in research protocols. 59 The invasive aspect of this technique with cannulation of peripheral arteries poses the potential for injury to the artery and nerves, which makes it less desirable for routine clinical

use and examination of larger populations. It is a valuable research tool to evaluate the pathological mechanisms underlying endothelial dysfunction and the impact of various therapeutic interventions.

The endothelium-dependent vasodilator response may Noninvasive Assessment of Peripheral Conduit Vascular Reactivity: Brachial Artery Ultrasound

Ultrasound techniques have long been used to study vessel physiology. Ultrasound assessment of brachial artery vasore-activity with high-resolution B-mode ultrasound emerged as a clinical substance P, or to a physical stimulus, such as shear stress research tool in the early 1990s to noninvasively study endotheliumdependent vasomotor function. In 1992, Celemajer and colleagues used B-mode ultrasound to study the brachial artery's vasoactive response to increased shear stress induced by hyperemia in adults with coronary artery disease or smoking exposure. 52,53 The key observation, termed flow-mediated vasodilation (FMD), expressed as a percentage (FMD%), was impaired vasoactivity detectable noninvasively with ultrasound in individuals with risk factors, coronary artery disease, or both. Concurrent with observations from basic studies and invasive coronary studies, flow-mediated vasodilation in the brachial artery is abolished by the NO synthase inhibitor 1-NMMA. 60,61

Since its inception, flow-mediated vasodilation has been widely applied as a research tool to evaluate the impact of cardiovascular risk factors and preclinical disease states and to improve endothelial function with targeted specific interventions and risk factor modifications. Further exploration of the technique led to studies of underlying mechanisms for the time course after the hyperemic response, the effect of blood pressure cuff occlusion duration, the assessed by measurement of coronary blood flow with effect of upper arm occlusion versus lower arm occlusion for intravascular Doppler study in response to intra coronary technical ease of imaging and targeted NO stimulation, and the effect of other stimuli such as cold pressor and mental stress. 62-69

Advances in vascular biology and the role of oxidative stress in enhancing lipid oxidation and promoting a pro-inflammatory state spearheaded further investigations with this technique. Basic studies of vascular rings exposed to chylo micron remnants 70 and the effect of fat meals with and without antioxidant vitamins on flowmediated vasodilation 71,72 contributed to the notion that healthy vascular beds exposed to oxidative stressors manifest acute alterations in endothelial function. Studies that demonstrated significant improvement in flow-mediated brachial artery vasodilation inspired enthusiasm to study the impact of various risk factors and targeted therapies for cardiovascular disease. Correlative studies of brachial artery vasoactivity with intra coronary vasoactive substances and carotid intima-media thickness provided support for brachial artery vasoactivity as an index or marker of endothelial function. 73,74 Large clinical trials and epidemiological longitudinal studies now incorporate FMD% rate as a marker of impaired vasomotor endothelial dysfunction.

interest to determine whether there were methods to more vasoactivity testing are well described in the literature. Seemingly simple, it is technically challenging, with a significant learning curve to achieve high-quality, consistent performance and reproducibility in both technique and interpretation. 63-65,74,75

Standard ultrasound systems equipped with vascular software relationship of conduit and resistance vessels in vascular with B-mode two-dimensional imaging, color and spectral Doppler display, internal electrocardiographic monitor, and high-frequency vascular linear array transducer (range, 8 to 12 MHz) are required. technique that indirectly measures microvessel function as Higher frequency transducers of 8 to 12 MHz compared with 7 to 7.5

vasoactive substances or stimuli and rest quietly supine for 10 after hyperemia for off-line analysis (Fig. 32-3). minutes.

extended and supinated to allow optimal imaging of the brachial artery antecubitally. Blood pressure occlusion for 5 minutes and release create the shear stress to induce flow-mediated vasodilation.

Application of an upper versus a lower arm cuff for occlusion has been studied, and the placement provides different stimuli. Upper arm occlusion provides a more robust shear stimulus for vasodilation and probably invokes other vasoactive mechanisms hyperemic blood flow are calculated from the time-averaged pulsed increase in hyperemic flow of four to five times from baseline and a of one waveform to the beginning of the next waveform. greater change in vasodilation compared with lower arm cuff Blood flow velocity (TVI) x vessel diameter (n r 2) occlusion. Furthermore, vasodila tor response to upper arm occlusion hyperemia has been shown to be largely NO mediated. In addition, arteries smaller than 2.5 mm in diameter are difficult to

measure, and vaso dilation is difficult to perceive in vessels larger than 5.0 mm in diameter.

Flow-mediated vasodilation typically occurs 60 to 90 seconds after cuff release while accurately acquiring the image at the same position compared with baseline. Brachial artery flow returns to high dose (0.4 mg) of nitroglycerin spray or sublingual tablet, has a single high dose (0.4 mg) of nitroglycerin spray or sublingual tablet, has a single high dose (0.4 mg) of nitroglycerin spray or sublingual tablet, has a single high dose (0.4 mg) of nitroglycerin spray or sublingual tablet, has a single high dose (0.4 mg) of nitroglycerin spray or sublingual tablet, has a single high dose (0.4 mg) of nitroglycerin spray or sublingual tablet, has a single high dose (0.4 mg) of nitroglycerin spray or sublingual tablet, has a single high dose (0.4 mg) of nitroglycerin spray or sublingual tablet, has a single high dose (0.4 mg) of nitroglycerin spray or sublingual tablet, has a single high dose (0.4 mg) of nitroglycerin spray or sublingual tablet, has a single high dose (0.4 mg) of nitroglycerin spray or sublingual tablet, has a single high dose (0.4 mg) of nitroglycerin spray or sublingual tablet, has a single high dose (0.4 mg) of nitroglycerin spray or sublingual tablet, has a single high dose (0.4 mg) of nitroglycerin spray or sublingual tablet, has a single high dose (0.4 mg) of nitroglycerin spray or sublingual tablet, has a single high dose (0.4 mg) of nitroglycerin spray or sublingual tablet, has a single high dose (0.4 mg) of nitroglycerin spray or sublingual tablet, has a single high dose (0.4 mg) of nitroglycerin spray or sublingual tablet, has a single high dose (0.4 mg) of nitroglycerin spray or sublingual tablet, has a single high dose (0.4 mg) of nitroglycerin spray or sublingual tablet, has a single high dose (0.4 mg) of nitroglycerin spray or sublingual tablet, has a single high dose (0.4 mg) of nitroglycerin spray or sublingual tablet, has a single high dose (0.4 mg) of nitroglycerin spray or sublingual tablet measured perpendicular to the longitudinal axis in which the lumen- increase from baseline diameter. Brachial artery images and interface is identified with electronic calipers from the ultrasound 30 minutes. system analysis software or determined through edge detection computer software typically performed off-line. Edge detection programs that can account for skew by elliptical modeling have less variance in their measurements of brachial artery diameter. 76 Maximal vasodilation after hyperemia may be variable within an individual or in certain disease states, although most individuals

of the intima as opposed to the vessel wall alone, which has dilate maximally at approximately 60 to 90 seconds after cuff release. implications for the application of measurement techniques. Before Consequently, it is recommended that B-mode acquisition of the brachial artery vasoactivity testing, the individual should avoid brachial artery be acquired from 30 seconds up to at least 2 minutes

Flow-Mediated Vasodilation. FMD% is expressed as the change The technical preparation involves positioning the subject 's arm in post-stimulus diameter as a percentage of the baseline diameter.

> FMD% = post - hyperemic diameter - baseline diameter + baseline **diameter** x 100

Doppler: Baseline and Hyperemic Flow. Baseline and besides release of NO. Upper arm occlusion typically elicits an Doppler spectral trace (TVI, time-velocity integral) from the onset32

x heart rate = blood flow

baseline by 1 minute after hyperemia, but dilation may ensue beyond been given to determine the maximum obtainable vasodilator 1 minute. ^{62,63} The physiological characteristics of this response may response and to serve as a measure of endothelium independent be individually dependent as well as indicative of impaired vascular vasodilation reflecting vascular smooth muscle function. ^{67,68} Peak health in the setting of risk factors or genetic predisposition to vasodilation occurs 3 to 4 minutes after nitroglycerin disease states such as atherosclerosis. Brachial artery diameter is administration, on the order of 15% to 20% maximum dilation intima interface is visualized on the near (anterior) and far Doppler velocity signal are continuously recorded during this time. (posterior) walls. The lumen-intima interface or media-adventitia The effect of nitroglycerin on the vessel diameter persists up to 20 to

TIME COURSE OF FLOW-MEDIATED DILATION (FMD) EXAM PROTOCOL

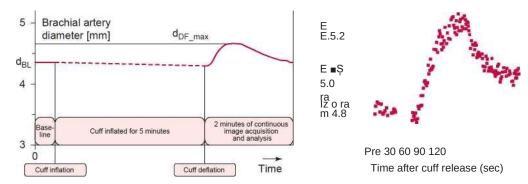


FIGURE 32-3 A, Schematic diagram of the brachial artery B-mode ultrasound imaging protocol for brachial artery vasoactivity testing. B-mode longitudinal images of the brachial artery segment of interest are recorded during baseline and after cuff deflation. Baseline and deflation image sequences are analyzed to assess vessel diameter function. Diameter d is the averaged arterial diameter at baseline; duration of baseline acquisition typically ranges between 10 and 20 seconds; d is the maximum diameter during the 2 minutes after cuff release. B, Time course of brachial artery flow-mediated vasodilation by upper arm occlusion shear stress stimulus in a healthy individual. (A from Sonka M, Liang W, Lauer RM: Automated analysis of brachial ultrasound image sequences: early detection of cardiovascular disease via surrogates of endothelial function. IEEE Trans Med Imaging 21:1271, 2002. B from Corretti MC, Anderson TJ, Benjamin EJ, et al: Guidelines for the ultrasound assessment of endothelial-dependent flow-mediated vasodilation of the brachial artery. J Am Coll Cardiol 39:257, 2002.)

534 Nitroglycerin should not be administered to individuals with clinically significant bradycardia or hypotension. Determination of the vasodilator responses to increasing doses of nitroglycerin (an NO donor), rather than to a single dose, may further elucidate changes in smooth muscle function or arterial compliance that might be playing a role in any observed changes in flow-mediated vasodilation. 67,68 This may further the understanding of impaired vasodilator responses in the presence of cardiovascular risk factors or disease states.

Quality Control Metrics and Reproducibility

32 Quality control of measures of flow-mediated vasodilation requires optimal hands-on training to ensure the highest quality and consistency in data acquisition and measurement techniques. 63 Intraobserver and interobserver variability for technical and interpretive expertise is highly dependent on the mastery of this technique. This requires a learning curve based on appropriate instruction, consistent application, and quality control. The image analysis and measurement of the vasodilator response from repeated studies should be done by an individual who is blinded as to sequence. Investigatorinitiated studies, large-cohort epidemiological studies, and core laboratories should adhere to a standardized protocol for training, implementation of technique and analysis, and reproducibility. Experienced laboratories with excellent reproducibility maintain a coefficient of variation < 2% and a 2% to 4% improvement in FMD% in small crossover trials (N = 20 to 30 subjects) and parallel group trials (N = 25 to 50) per intervention arm. 63,69,77,78

Interpretation

To date, there are no specific nomograms for flow-mediated vasodilation other than data from ongoing data bases such as the Framingham study or other individual institutions. necessary, along with consideration of physiological factors that mediate basal vasomotor tone and diameter. A larger known as pulse volume amplitude). baseline diameter yields a smaller measure of percentage change for any given absolute change in post-stimulus diameter. Conversely, smaller arteries dilate relatively more attempting to compare vasodilator responses between individuals and groups with different baseline diameters. For an intervention in the same individuals, percentage change percentage change in diameter are therefore typically reported in publications.

Flow-mediated vasodilation is also affected by a change in the hyperemic stimulus. The flow stimulus should be artery may be related to changes in flow (even indirectly Further studies with noninvasive techniques of vasomotion and of hyperemic flows as shear stimulus and its physiological significance are currently under way. 79-83

EMERGING TECHNIQUES OF VASCULAR

FUNCTION recordings with sphygmography, later replaced by the sphygmomanometer. Pulse characteristics and arterial hemodynamics have always been an important focus of the study of vessel function and ventriculoarterial coupling. Advances in vascular biology coupled with computer technology and techniques to measure arterial waveforms, pressures, and flows generated a greater interest in noninvasive study of vascular structure and function with age, other cardiovascular risk factors, and more recently the impact of disease and risk factor modification. The association of arterial stiffening and aging is well described.

Elastin and collagen, major components of the vasculature, largely regulate vascular tone or stiffness and undergo significant degenerative changes with age and other cardio vascular risk factors or disease processes, as in hypertension, diabetes, end-stage renal disease, and smokers. Arterial stiffness has been extensively studied and accepted as a surrogate marker of atherosclerosis and the effects of advancing age and as a prognostic indicator for cardiovascular -

Arterial stiffness is structurally determined by the components of the blood vessel, but it is partly under the functional control of the endothelium through its release of vasoactive mediators. The emerging data that arterial stiffness may be involved in the pathogenesis of cardiovascular disease underscore the interest in and importance of physiological mechanisms that may be targeted for risk assessment and therapeutics. Various techniques to noninvasively study vascular properties and function through assessment of arterial waveforms, pulse velocity, and measures of arterial stiffness, along with flow-mediated vasoactive surrogates of endothelial function, may be used selectively and comprehensively to fully evaluate the status of vascular function in an individual and the impact of risk factors and interventions. These include Dopplerbased techniques and applanation tonome try to assess velocities or Consistency in technical approach and data acquisition is flow and arterial waveforms include pulse wave analysis, pulse wave velocity measurement, and pulse amplitude tonometry (also

Given the challenges in mastering ultrasound assessment of brachial artery vasodilation in response to reactive hyperemia, interest has shifted to the utility of measuring digital pulse than larger arteries do. In sum, this merits consideration in amplitude tonometry at rest and in response to reactive hyperemia to assess cardiovascular risk in individual subjects.84

Pulse amplitude tonometry (commercially available as Endostudies in which multiple measures are made within the same PAT2000, Itamar Medical) is a relatively new non-invasive technique individual over time or comparisons are made before and after that records pulse amplitude in the fingertip at baseline and during reactive hyperemia. Hyperemia flow-mediated dilation in the may be the easiest metric, provided baseline diameter remains fingertip increases the pulse amplitude. With use of proprietary stable over time. The baseline diameter, absolute change, and software, the net response is expressed as the reactive hyperemia pulse amplitude tonomy index. Preliminary studies with this technique have demonstrated that an intra-arterial infusion of L-NMMA into the brachial artery blocks hyperemia-induced increase in pulse amplitude. This indicates that the reactive hyperemia pulse consistent; otherwise, any change in FMD% of the conduit amplitude is at least partially dependent on an NO synthesis process. 84-87 Other investigations have shown a correlation between reactive mediated by changes in the microcirculation) rather than to hyperemia-induced pulse amplitude tonometry index and brachial improve ment of endothelial function of the conduit vessel. artery flow-mediated vasodilation and the coronary circulation 85 and that it too is inversely related to cardiovascular risk factors, 85,86 as previously shown with brachial artery flow-mediated vasodilation and other invasive techniques.

> The technique can be simultaneously employed with ultrasound assessment of brachial artery flow-mediated vaso dilation to provide further insights into hyperemia-induced vasomotor function. Pulse amplitude tonometry appears to be a promising technique, particularly given its ease of application; however, more studies are

needed to confirm

its correlation with other well-established techniques and protocols. aorta and carotid, brachial, or femoral arteries.

Microvessels

An emerging noninvasive technique to study the microvasculature is laser Doppler flowmetry, which provides a semi-quantitative measure of blood flow in the small blood vessels of the microvasculature. These vessels have low-velocity flows associated with nutrient blood flow delivery, regulation of skin temperature, and vascular resistance in the capillaries, arterioles, and venules. A low-intensity laser beam scans across the skin surface, allowing measurements with good temporal and spatial resolution of rapid blood flow changes. Application of this technique to the study of endothelium dependent and endothelium-independent vasomotor function with use of substances that change skin perfusion, such as acetylcholine, is under study; therefore, its exact role remains uncertain.

Arterial Compliance and Arterial Stiffness

Each pulsation of the heart generates a velocity of the pressure wave (pulse wave velocity) that is transmitted centrally throughout the peripheral vascular system. The waveform is related to the biomechanical properties of the arterial system, including arterial wall stiffness. The ascending aortic pressure waveform can be measured from the carotid artery, femoral artery, or radial artery by noninvasive techniques such as applanation tonometry and Doppler ultra sound. Arterial stiffness can be examined with a variety of approaches. These include measurement of the arterial pulse pressure from blood pressure readings, applanation tonome try assessment and analysis of the pulse wave contour, and ultrasound assessment of the arterial distensibility and pulse wave velocity. 88,89

Pulse wave analysis is a technique that provides accurate recording of peripheral pressure waveforms with assessment of the corresponding central waveform. From this, the augmentation index and central pressure can be derived, providing information about arterial stiffness and altered pulse dynamics. Pulse wave analysis is a stable, easy to perform, and highly reproducible technique that relates arterial structure to vascular tone. Implementation of the technique is not dependent on a flow stimulus or infusion of a vasodilator agent. Pulse wave analysis is typically applied to the

Applanation tonometry (SphygmoCor CPV, AtCor Medical, Sydney, Australia) is a method used for pulse wave analysis to Laser Doppler Flowmetry: Measure for Cutaneous derive the central aortic pressure waveform and pulse wave velocity. There are two main methods of applanation tonometry to assess the pulse pressure waveform. It can be analyzed by applying a valid transfer function based on Fourier analysis, which allows derivation of an aortic pressure waveform, the augmentation index, and time to reflected wave. The other method involves the use of a modified windkessel model of the circulation. This model allows calculation of large- and small-vessel compliance but does not include the effects of wave reflections. The tonometer is positioned over the maximal arterial pulsation of an accessible superficial artery (eg, radial, brachial, and femoral) to minimally flatten or applanate the 32 arterial wall. This results in normalization of the circumferential stress in the arterial wall. Changes in the electrical resistance of a piezoelectric crystal within the tonometer allow recording of the pressure wave form. This pressure waveform is then digitized for application of a "generalized transfer function," derived from simultaneous recording of peripheral arterial and the invasively recorded central ascending aortic pressures, allowing automated calculation of the central aortic pressure waveform from the peripheral pressure and tonometry data. Reflection of pressure waves in the arterial tree leads to augmentation of the central aortic pressure wave. Peripheral vessels with higher impedance reflect the 🭠 incident wave to the aorta, resulting in augmentation of the central aortic pressure. Thus, the resulting central aortic pressure represents the sum of the incident and the reflected waveforms. 90

The obtained pulse waveform shape provides information about arterial compliance and serves as the basis for the calculation of the augmentation index as illustrated in Figure 32-4. Augmentation index is commonly used as a measure of arterial stiffness. Applanation tonometry-based assessment of peripheral pressure waveforms and calculation of augmentation index have been studied as a tool for assessment of endothelial function. However, in a comparison between flow-mediated vasodilation and pulse wave analysis as a

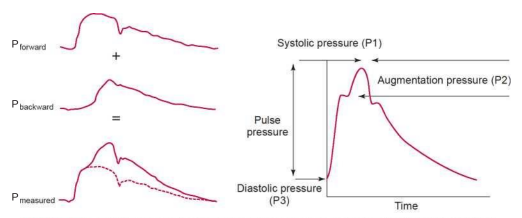


FIGURE 32-4 Left. Diagram of agric pressure wave. Invasive hemodynamic measure of agric pressure waveforms: forward, backward, and summated to yield measured pressure and flow waveforms. Right. Applanation tonometry: pulse wave analysis and augmentation index assessment of arterial stiffness. Augmentation index = augmentation pressure/ pulse pressure. The peak systolic pressure is represented by P1. P3 is the minimum diastolic pressure. An inflection point, P2, in the waveform identifies the merging point of the beginning upstroke of the reflected pressure wave. (Left from Murgo JP, Westerhof N, Giolma JP, Altobelli SA: Manipulation of ascending aortic pressure and flow wave reflections with the Valsalva maneuver: relationship to input impedance, Circulation 63:122, 1981. Right from Laurent S, Cockcroft J, Van Bortel L, et al: Expert consensus document on arterial stiffness: methodological issues and clinical applications, Eur Heart J 27:2588, 2006.)

536 function of endothelial function in response to inflammation, flow-mediated vasodilation was found to be more reproducible. Multiple variables (age, height, and heart rate) may influence calculations. In addition, standardization of arteries used for calculations needs to be determined. Paucity of long term longitudinal studies assessing the predictive value of these tests on mortality and the effect of interventions is a limitation in wide-scale adoption of this tool.

Augmentation index and pulse wave velocity are dependent on arterial stiffness and ventricular-arterial coupling and I wave reflections that include effects of the mechanical properties of the arterial wall in addition to the effects of atherosclerosis and dysfunctional endothelium. This is particularly notable in renal failure, hypertension, presence of advanced 32 glycation end products in diabetes mellitus, and advanced age. There may be discordance with FMD% and some of these other techniques, which speaks to the additive value of the two methods.

LIFESTYLE MODIFICATIONS AND THERAPEUTICS

contributor to cardiovascular and in diseases such as diabetes, dysmetabolic syndrome, role in atherosclerosis. This small group of studies hypertension, and coronary artery disease. Endothelial dysfunction also invokes the inability to optimally repair in the

setting of vascular injury. The presence of endothelial dysfunction represents in itself a risk factor burden that seeks to be assessed and quantified; however, the complexity and multifactorial nature of endothelial dysfunction eludes simple diagnostic tests. It is the final common pathway through which vascular risk factors initiate the pathogenesis of atherosclerosis. Current evidence supports the fact that endothelial function is an integrative indicator of the net effects of the magnitude and duration of arterial injury and repair over time. The focus on specific disturbances of endothelial dysfunction allows the application of noninvasive imaging and hemodynamic techniques with or without concurrent biomarkers. Numerous studies employing flowmediated vasodilation as a marker of endothelial vasomotor dysfunction have demonstrated impairment (less vasodilation in response to shear stress) in various conditions and improvement with such interventions as life style modifications and therapeutics.

FORECAST

Numerous studies have emerged in the last decade that employed There is a growing body of evidence that endothelial several techniques to measure endothelial dysfunction as a potential dysfunction can be both a consequence of and a major surrogate marker for clinical events. Table 32-3 summarizes a variety disease and events of retrospective and prospective studies of endothelial function Experimental and clinical techniques have demonstrated testing by invasive and noninvasive modalities with cardiovascular various manifestations of endothelial dysfunction in the events. 94-103 These studies lend further support to other experimental presence of traditional and novel cardiovascular risk factors and clinical evidence that the endothelium plays an active critical -

Author	Design and Patient Cohort	Vascular Bed	Technique	Stimulus	Follow-up	Cardiovascular Events
Suwaidi (2000)	Retrospective 157 mild CAD	Coronary	IVUS, QCA/CBF	Acetylcholine adenosine, nitroglycerin	28 months	14%, 10 events Events with lowest acetylcholine response
Schachinger (2000) Re	etrospective 147 CAD	Coronary	IVUS, QCA/CBF	Acetylcholine, cold	7.7 years	11%, 28 events Acetylcholine independent predictor of events
Neunteufl (2000)	Retrospective 73 CAD	Brachial	FMD%	Hyperemia	5 years	FMD% independent predictor of events
Heitzer (2001)	Prospective 281 CAD	Brachial	Venous plethysmography	Acetylcholine	4.5 years	Acetylcholine independent predictor of events
Perticone (2001)	Prospective 225 HTN	Brachial	Venous plethysmography	Acetylcholine	32 months	Acetylcholine independent predictor of events
Gokce (2002)	Prospective 187 preoperative vascular	Brachial	FMD%	Hyperemia	30 days	15% (FMD% < 8%) FMD% independent predictor o postoperative events
Halcox (2002)	Retrospective 308 diagnostic catheterization	Coronary	QCA/CBF	Acetylcholine, adenosine	46 months	11.4% Acetylcholine independent predictor of events
Modena (2002)	Prospective 400 postmenopausal women with HTN	Brachial	FMD%	Hyperemia	67 months	21.3% with persistent impaired FMD%; no improvement with therapy
Schindler (2003)	Prospective 130 normal coronary angiograms	Coronary	Vasoactivity	Cold pressor	45 months	26 with events Cold pressor independent predictor of events
Gokce (2003)	Prospective 199 vascular surgery	Brachial	FMD%	Hyperemia	14 months	Cardiovascular death, unstable angina, stroke

CAD, coronary artery disease; CBF, coronary blood flow; FMD, flow-mediated vasodilation; HTN, hypertension; IVUS, intravascular ultrasound; QCA, quantitative coronary angiography

demonstrates prognostic insight into selected high-risk groups with or in addition to traditional clinical tools, particularly for subclinical disease demonstrate a wide range of responses to endothelium - in the identification of individuals who might benefit from various dependent techniques and heterogeneity in the magnitude of thera pies. An improvement displayed by endothelial function dysfunction in individuals with similar risk factor profiles. With testing should predict a measurable reduction in the risk of clinical advances in development and application, such techniques will cardiovascular events. Genetic susceptibility as well as complex serve as useful diagnostic and prognostic clinical and research tools. multiple risk factor interactions throughout the continuum of

CLINICAL TRIALS

The study of endothelial function in clinical research has had a universal interest as an important functional barometer of cardiovascular risk as well as a marker of endothelial injury coupled with other circulating biomarkers and emerging imaging techniques. Throughout the 1990s, pharmacological and lifestyle modifications known to decrease cardiovascular risk were studied with endothelial function testing as a clinical endpoint. Numerous clinical studies reported the effect of pharmacologic or physiologic interventions on endothelial function by brachial artery flow-mediated vasodilation and other endothelial function in healthy individuals and in altered endothelial vasomotor function in healthy individuals and in studies, both parallel group and crossover designs, were successfully Understanding of the pathophysiology of endothelial dysfunction conducted on the effects of angiotensin-converting enzyme due to underlying injury and repair has made significant progress. literature is variable in this regard.

and sample size determinations have been reported. Subsequently, and re-endothelialization. assessment of flow-mediated brachial artery vasodilation has been analysis.

To date, there is no direct evidence that therapeutic improvement in endothelial function translates into lower cardiovascular be they imaging modalities of structure and function along with morbidity and mortality because large prospective clinical trials with circulating biomarkers - will provide improved endothelial function as a primary therapeutic endpoint have not been conducted. This is in part due to the technical and physiological challenges in optimizing standardization and reproducibility consistently to study the dynamic nature of the endothelium. Further development and investigation of various non-invasive techniques to clinically assess the functional properties of normal and activated endothelium are essential as no one technique can be used exclusively to assess and to monitor the dynamic nature of vascular biology. Currently, endothelial function testing is not used clinically to routinely assess risk or to guide management of risk factor modification or therapeutics. At this time, these methods have yet to emerge as powerful diagnostic tools alone

multiple risk factors for atherosclerosis that have an impact on atherosclerosis and vascular disease. Ongoing studies with these endothelial function; data to support the prognostic value in low- techniques continue to address their full potential. This requires and intermediate-risk groups are currently lacking. Nevertheless, endothelial function testing to be sensitive, specific, and accurate. It these prognostic studies suggest that endothelial function testing should be standardized and reproducible along with validated may provide insight into an individual's risk burden, the response to population norms to help provide insight into the status of vascular therapy, and perhaps the need for more aggressive management. All health and possibly guide therapy. It should provide prognostic biological systems, individuals with cardiovascular risk factors, and information to current clinical information and parameters and assist vascular health and disease will likely determine any one32 individual's response to testing. The complex causal mechanisms of atherosclerosis, the diverse effects of various interventions imposed along the continuum of vascular health, and the genetic-environmental response to such interventions make it unlikely that any single surrogate of endothelial function can optimally stratify

inhibitors, 52,53,57 antioxidant vitamins, 52,57 statins, 52,57,69 diet changes, Whereas risk factors associated with endothelial dysfunction are ^{69,71} and hormone replacement therapy ^{52,69} on endothelial function, increasingly recognized, detection of endothelial dysfunction is still putatively through improvement in NO-mediated mechanisms, primarily a research tool. With advances in biomarkers and imaging Although several intervention therapies demonstrate improvement technology, it may be possible to detect injury early in the process in endothelial function and also reduce cardiovascular risk, the before expression of the cardiovascular phenotype. In addition, with recent advances in the repair process, it will be imperative to identify Implications of the approach for standardization of techniques - and track circulating endothelial progenitor cells to document repair

Various invasive and noninvasive techniques have been applied incorporated into clinical studies and larger longitudinal studies to further study the endothelial vasomotor dysfunction in the wake such as the Framingham study and other similar epidemiological- of the discoveries made in NO biology and oxidative injury. genetic studies. There are technical and interpretive challenges that Ultrasound assessment of brachial artery flow-mediated must be mastered to ensure consistency and reproducibility in data vasodilation has yielded important information about vascular acquisition owing to the variable nature of vascular reactivity. function in health and disease, yet several new technological Several important considerations are optimal technical training, advances and techniques have emerged to expand the scope in the study design, and sample size along with uniform technique in detection of mechanisms of and impairments in vascular structure scanning protocol and analysis and validation of reproducibility in and function. The ongoing discoveries from basic experimentation to controlled research settings. Most studies have been of cohorts of translational clinical studies remain essential to the study of individuals with risk factors or disease states that undergo mechanisms of impaired vascular homeostasis and to determine the assessment of endothelial function testing for diagnostic or increased mental value of the presence of endothelial dysfunction therapeutic assessment from single centers and less so from beyond known and established risk factors and clinical predictors. multicenter studies. Multicenter studies opti mally require one site Whether treatment strategies aimed at modifying risk factors are serving as the core laboratory to ensure uniform methodology and sufficient or a need for endothelium-directed therapy is required is an exciting area of investigation and needs to be proven.

Diagnostic and prognostic studies of endothelial dysfunctions –

538 insight and guidance into an individual's overall cardiovascular risk burden, the treatment plan, and the response to therapy. These goals for brachial artery vasoactivity testing and other related noninvasive vascular techniques require rigorous standardization 33. and correlations, data from large-scale population and randomized studies, and correlation studies with other imaging modalities coupled with genomics and environmental research. Prospective studies comparing vascular function testing with modalities such as intravascular ultrasound, carotid intima-media multidetector computed tomography, and computed tomographic angiogra phy in the context of interventions and prognosis will further define the potential clinical role for ultrasound measures of 38. Mel LG, Pachori AS, Kong D, et al: Endothelium-targeted gene and cell-based therapy for brachial artery vasoactivity testing with other emerging techniques and biomarkers to assess cardiovascular health and disease.

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SECTION IX

Exercise/Emotional Aspects of Preventive Cardiology

CHAPTER 33

Exercise for Restoring Health and Preventing Vascular Disease

Kerry J. Stewart, Elizabeth V. Ratchford, and Mark A. Williams

- Elderly individuals with heart disease can benefit greatly from exercise training and other aspects of cardiac rehabilitation and secondary prevention programs.
- Exercise training plays a critical role as a primary treatment of patients with peripheral arterial disease, with the goal of improving quality of life and functional capacity.

KEY POINTS

factors, metabolic syndrome, and diabetes.

| Increased levels of physical diabetes activity and exercise are associated with increased

- By adding a sedentary lifestyle to crease in its list of controllable risk factors cardiovascular disease risk for coronary artery disease, the American Heart Association has made regular exercise a major focus for preventive medicine.
- Exercise regimens should include aerobic, muscle strengthening, and flexibility exercises.
- Cardiac rehabilitation is recognized as integral to the comprehensive care of patients with cardiovascular disease and is recommended as useful and effective (Class I) by the American Heart Association and the American College of Cardiology in patients with coronary artery disease and chronic heart failure.
- The efficacy and effectiveness of cardiac rehabilitation on improvements in health outcomes are beyond the improvements in morbidity and mortality already available through revascularization and optimal pharmacotherapy.

This chapter describes the cardiovascular health benefits of regular exercise, the benefits and risks of exercise training and cardiac rehabilitation for individuals with established cardiovascular disease including peripheral arterial disease, and the major types of exercise recommended for cardiovascular health. Exercise prescription guidelines are provided to ensure maximal efficacy and safety of the exercise program.

ROLE OF INCREASED PHYSICAL ACTIVITY IN PRIMARY PREVENTION OF CARDIOVASCULAR DISEASE

The cardiovascular health benefits of regular exercise are well established. ¹⁻³ According to the US Department of Health and Human Services, ³ strong evidence demonstrates that compared with less active persons, more active men and women have lower rates of all-cause mortality, coronary heart disease, hypertension, stroke, type 2 diabetes, metabolic syndrome, colon cancer, breast cancer, and depression. The risk of dying prematurely declines as people become physically active, as evidenced by the data shown in Table 33-1. ¹⁻³

Strong evidence also supports the conclusion that compared with less active people, physically active adults and older adults exhibit a higher level of cardiorespiratory and muscular fitness, have a healthy-body mass and composition, and have a metabolic profile that is more favorable for prevention of cardiovascular disease and type 2 diabetes. Modest evidence indicates that physically active adults and older adults have better quality sleep, health-related quality of life, and enhanced bone health.

A recent meta-analysis 4 of 26 studies, incorporating 513,472 individuals (20,666 coronary heart disease events) followed up for 4 to 25 years, showed that individuals who reported performing a high level of leisuretime physical activity had significant protection against coronary heart disease (relative risk, 0.73; P < 0.00001), whereas those individuals who practiced a moderate level of physical activity also had a reduced risk of coronary heart disease (relative risk, 0.88; P < 0.0001). Data from the Physicians' Health Study 5 showed that adherence to healthy lifestyle factors (normal body weight, never smoking, regular exercise, moderate alcohol intake, and consumption of breakfast cereal and fruits and vegetables) is independently

Minutes per Week of Moderate- or	ntensity Physical Activity Relative Risk 1 0.8 0.73
Vigorous-Intensity Physical Activity	Relative Risk
30	1
90	0.8
180	0.73
330	0.64
420	0.615

associated with an 11% lifetime risk of heart failure with exercising > 5 times per week versus a 14% risk for exercising < 5 times per week. Thus, there is substantial protection against the occurrence of cardiovascular disease from moderate to high levels of physical activity, thereby strengthening the recommendations of guidelines. 6

Increased levels of physical activity and exercise are also associated with increased longevity and a decrease in cardio vascular disease risk factors, metabolic syndrome, and diabetes. In 2009, the American College of Sports Medicine's position statement "Exercise and Physical Activity for Older Adults" 7 reported that although no amount of physical activity can stop the biological aging process, there is evidence that regular exercise can minimize the physiological effects of an otherwise sedentary lifestyle and increase active life expectancy by limiting the development and progression of chronic disease and disabling conditions. There is also emerging evidence for significant psychological and cognitive benefits accruing from regular exercise participation by older adults.

Ideally, exercise prescriptions for older adults should include aerobic exercise, muscle strengthening exercises, and flexibility exercises. Although this chapter focuses on cardio vascular disease, adults with chronic conditions such as type 2 diabetes, cancer, osteoarthritis, cognitive disorders, and renal disease are also likely to derive significant therapeutic benefit from prescribed exercise and increased physical activity. 3 In many cases, persons with cardiovascular disease also suffer from these comorbidities, thereby enhancing the importance of exercise training as part of their overall treatment plan.

Despite numerous epidemiological and experimental studies supporting the beneficial effects of being physically active, this scientific knowledge has not resulted in a more active population. According to the Centers for Disease Control and Prevention, 8 the prevalence of leisure-time physical inactivity throughout the United States ranged from 16% to 49%, whereas the prevalence of adults who engaged in at least moderate physical activity ranged from 33% to 62%, and the prevalence of vigorous physical activity ranged from 15% to 42%. Among adults aged 60 years or older, more than half reported no leisuretime physical activity at all. ⁹Thus, a substantial number of adults do not engage in levels of activity sufficient to produce health benefits.

People with chronic disease are more likely to report a sedentary lifestyle, as are minorities and individuals in lower socioeconomic classes. 10 Unfortunately, less than 30% of individuals at high risk for cardiovascular disease receive physical activity counseling during ambulatory care visits. ¹¹ By adding a sedentary lifestyle to its list of controllable risk factors for coronary artery disease, the American Heart Association has made regular exercise a major focus for preventive medicine. 6

The physical activity goal in the Healthy People 2010 report by the Centers for Disease Control and Prevention is to increase the proportion of adults who engage in regular exercise and moderate activity for at least 30 minutes each day. This goal matches the Surgeon General's recommendation that all Americans should accumulate at least 30 minutes of activity

throughout the day on most days of the week. 12 These guidelines establish the minimal effort needed to derive health benefits from physical activity, although individuals exceeding these recommendations in terms of frequency or intensity are likely to derive additional health and fitness benefits.

Several studies have established an inverse dose-response relationship between the amount of physical activity per formed and health risk. Thus, the most fit and active individuals generally have the best risk profiles and reduced levels of early mortality and morbidity from a variety of diseases, including cardiovascular disease. For example, men and women who were generally healthy but less fit by exercise testing had a higher risk of all-cause mortality during an 8-year follow-up compared with those who were moderately or highly fit. Data from the Nurses' Health Study ¹³ found that physical activity of less than 3.5 hours per week and excess body weight (defined as a body mass index of 25 or higher) accounted for 31% of all premature deaths, 59% of cardio vascular deaths, and 21% of cancer deaths.

In another study, Puerto Rican men in the middle of the physical activity distribution had a 32% to 37% reduction in risk for all-cause mortality. 14 Moreover, compared with the most sedentary quartile of participants, the next most active quartile had an accumulated survival that was approximately 3 years longer. We previously showed that exercise-induced reductions in total and abdominal obesity after 6 months of training were associated with favorable changes in risk factors for cardiovascular disease, including those that constitute metabolic syndrome. 15

TRAINING EFFECT EXERCISE

The underlying principles for enhanced cardiovascular and musculoskeletal function are similar for those with and without cardiovascular disease and also apply to most other chronic health conditions, such as type 2 diabetes. 16 A brief summary of these principles is provided.

The body adapts to the kind and amount of physical demands placed on it. The response is specific, with the most change generally occurring in those parts of the body on which demands are placed. For exercise to improve fitness, it must overload the muscles or organ system involved in the exercise. In this context, overload is defined as performing exercise at a greater intensity than the intensity to which one is accustomed. Because the effects of exercise are specific to the type of activity engaged, an optimal exercise program should include a variety of activities designed to improve each of the major components of fitness. These have cardiovascular endurance, muscle strength, muscle endurance, and flexibility.

Cardiovascular endurance depends on the ability of the muscles to use oxygen during exercise; hence, it is also known as aerobic endurance. Cardiovascular endurance is dependent on cardiac output providing sufficient oxygen carrying blood, distribution of the blood to the working muscles, and capacity of the muscles to extract oxygen from a given blood flow.

The main hemodynamic adaptation to aerobic exercise in individuals with and without heart disease takes place in the peripheral vascular and muscular systems. With regular exercise, the skeletal muscles can extract more oxygen from a given blood flow, and there is a better distribution of the cardiac output. Heart rate and blood pressure are also lower at rest and at a given submaximal workload. As a result, the individual can do more work with less cardiac effort. This

adaptation is especially beneficial to cardiac patients who may have limited coronary artery blood supply or poor left ventricular function. Angina may occur at the same threshold, that is, the same double product (heart rate x systolic blood pressure), but this threshold is reached at a higher level of body work.

RISKS OF INCREASED PHYSICAL ACTIVITY

Although habitual physical activity reduces coronary heart disease events, vigorous activity can also acutely and transiently increase the risk of sudden cardiac death and acute myocardial infarction in susceptible persons, as discussed in detail in an American Heart Association Scientific Statement. ¹⁷ Exercise-associated acute cardiac events generally occur in individuals with structural cardiac disease. Hereditary or congenital cardiovascular abnormalities are predominantly responsible for cardiac events among young individuals, whereas atherosclerotic disease is primarily responsible for these events in adults.

The absolute rate of exercise-related sudden cardiac death varies with the prevalence of disease in the study population. The incidence of acute myocardial infarction and sudden death is greatest in the habitually least physically active individuals. Although no specific strategies have been widely studied to reduce exercise-related acute cardiovascular events, maintaining regular physical activity may help reduce events because a disproportionate number of events occur in the least physically active subjects performing unaccustomed physical activity. Other strategies, such as screening patients before participation in exercise, excluding high-risk patients from certain activities, promptly evaluating possible prodromal symptoms, training fitness personnel for emergencies, and encouraging patients to avoid high-risk activities, appear prudent but have not been systematically evaluated.

CARDIAC REHABILITATION

Cardiac rehabilitation and secondary prevention programs are recognized as integral to the comprehensive care of patients with cardiovascular disease and as such are recommended as useful and effective (Class I) by the American Heart Association and the American College of Cardiology in the treatment of patients with coronary artery disease and chronic heart failure. ¹⁸ In 2009, ¹⁹ an estimated 785,000 Americans had a new coronary event, and about 470,000 had a recurrent event. It is estimated that an additional 195,000 silent first myocardial infarctions occur each year. At the same time, the death rate from cardiovascular disease declined by 26% from 1995 to 2005. Thus, the burden of chronic car diovascular disease remains high, and a growing number of patients will be candidates for cardiac rehabilitation.

In 2007, 18 the American Heart Association and the American Association of Cardiovascular and Pulmonary Rehabilitation recommended that cardiac rehabilitation programs provide several important core components consisting of baseline patient assessment, nutritional counseling, risk factor management (lipids, hypertension, weight, diabetes, and smoking), psychosocial management, physical activity counseling, and exercise training. The American Heart Association also recommends these cardiac rehabilitation components for the elderly. 20 Whereas secondary prevention therapies, such as pharmacologic management of atherosclerosis risk factors and depression, are provided by clinicians in their offices, cardiac rehabilitation is often the most advantageous setting for bringing many of these core components together in a comprehensive approach to provide exercise training, patient education, behavioral counseling, and psychosocial support.

Studies of efficacy and effectiveness of cardiac rehabilitation

document reductions in mortality and improvements in clinical and behavioral outcomes beyond the improvements in morbidity and mortality already available through revas - cularization and optimal pharmacotherapy. Yet cardiac - rehabilitation, like many preventive measures, is underused. ²¹

Unfortunately, it is estimated that only 10% to 40% of eligible patients participate in cardiac rehabilitation programs. ^{22,23} These rates of participation are even less for older patients, a group with the highest prevalence of cardiovascular disease. ^{24,25} Although Medicare provides payment for cardiac rehabilitation for the diagnoses of myocardial infarction, coronary artery bypass surgery, stable angina, and, more recently, percutaneous revascularization, heart transplantation, and heart valve surgery, there is no such payment for patients who have heart failure or peripheral arterial disease. Insurance coverage by other third-party payers varies considerably throughout the United States.

There are also numerous disparities in cardiac rehabilitation program participation; women with lower incomes are less likely to be referred and less likely to enroll in cardiac rehabilitation, and there is a strong trend for African American women to be less likely to be referred and to enroll. ²⁶ Among 1933 cardiac patients who met the selection criteria of the American College of Cardiology guidelines of eligibility for cardiac rehabilitation, whites were more likely to be referred for cardiac rehabilitation than were blacks. ²⁷ Because almost all patients who have had an acute coronary event, with or without revascularization procedures, will benefit from cardiac rehabilitation, automatic referral systems should be considered to increase use and to reduce disparities. ²⁶

To compound the problem of access to cardiac rehabilitation based on race, blacks are more likely to have a greater number of adverse risk factors compared to whites. ²⁸ Although both groups gained secondary prevention benefits, the degree of improvement was less for blacks than for whites, and this was especially evident among black women.

A review of all components of cardiac rehabilitation and secondary prevention (such as smoking cessation, behavioral counseling, and pharmacotherapy) is beyond the scope of this chapter. To some extent, the scope is also limited by the nature of the studies on cardiac rehabilitation, which used supervised exercise training as the primary treatment modality. Although not fully evaluated in large-scale trials, a comprehensive approach to cardiac rehabilitation would be more than just exercise training and presumably would produce greater improvements in health and functional status for patients than is evident in the literature. ²⁹ Nevertheless, there is considerable evidence that exercise training by itself produces substantial physiological benefits, improves risk factors, reduces mortality, and increases aerobic capacity, muscle strength, and functional performance.

Regular exercise training, with the goal of achieving a training effect , is beneficial for almost all patients after myocardial infarction. ³⁰ Contemporary cardiac rehabilitation programs are also experienced in addressing the educational deficiencies of patients, including the special needs of those with cardiac pacemakers and implanted defibrillators, chronic heart failure, diabetes, peripheral arterial disease, and other comorbidities.

In recent years, there is also increased recognition of the importance of resistance training for individuals with and without cardiovascular disease. The American Heart Association recently updated its Scientific Advisory "Resistance Exercise in Individuals With and Without Cardiovascular Disease." ³¹ It notes that after appropriate screening, resistance training is an effective method to improve muscle strength

544 and endurance, to prevent and manage a variety of chronic medical through reductions in systemic inflammation. 35 conditions, to modify cardiac risk factors, and to enhance and use of elastic stretch bands.

studies show that cardiac patients who were required to carry or lift to prompt medical evaluation and treatment. weights or to perform isometric exercise after a myocardial infarction had fewer ischemic electrocardiographic changes and interventions, there are few controlled studies of cardiac arrhythmias during resistance exercise than during aerobic rehabilitation after these procedures. In one study, ³⁶ patients who

recreational activities require static efforts.

KEY BENEFITS OF EXERCISE TRAINING IN PATIENTS WITH CARDIOVASCULAR DISEASE

Coronary Artery Disease

Candidates for cardiac rehabilitation services historically were patients who recently had a myocardial infarction or had undergone coronary artery bypass graft surgery but now include patients who have undergone percutaneous coronary interventions, are heart transplantation candidates or recipients, or have stable heart failure, peripheral arterial disease with claudication, or other forms of other cardiac surgical procedures, such as those with valvular heart myocardial infarction. 42 disease, also benefit from such programs. 32

Patients who exercise regularly increase their physical working capacity and are therefore able to perform at higher levels of effort with less fatigue. Among those patients who experience exertional angina pectoris, regular exercise leads to cardiovascular adaptations as described earlier that result in the occurrence of angina at higher exercise levels. This increased anginal threshold allows the patient to do more work, and at any given level of work, the patient feels more comfortable because the work represents a lower percentage of a higher maximal capacity.

As summarized in the American Heart Association Scientific - Heart Failure Statement "Cardiac Rehabilitation and Secondary Prevention of Coronary Heart Disease," 30 exercise training, as part of a comprehensive rehabilitation program, has been shown to slow the progression or partially reduce the severity of coronary atherosclerosis. Multiple factors directly or indirectly appear to contribute to this effect. For example, increased flow-mediated shear stress on artery walls during exercise results in improved endothelial function, which is associated with enhanced synthesis, release, and duration of action of nitric oxide. Nitric oxide is responsible for endothelium-dependent vasodilation and inhibits multiple processes involved in atherogenesis and thrombosis.

Hambrecht and colleagues 33 demonstrated a significant improvement in endothelium-dependent arterial dilation in patients with coronary heart disease and abnormal endothelial function after only 4 weeks of vigorous endurance exercise training. Arterial inflammation probably plays a key role in the development and progression of atherosclerosis, as evidenced by the fact that acute myocardial infarctions often evolve from mild to moderate coronary artery stenoses and that patients who experience a fatal coronary event invariably had antecedent exposure to one or more major coronary risk factors. 34 Thus, another potential mechanism by which exercise training and increased cardiorespiratory fitness improve prognostic markers in persons with and without heart disease is

Percutaneous coronary interventions are effective for psychosocial well-being. Although weight machines are most interruption of the process of acute coronary stenosis. Although it is commonly used in formal cardiac rehabilitation programs, of great benefit that myocardial tissue damage can be avoided or alternative modes of resistance training are calisthenics, isometrics, minimized if the patient is treated in a timely manner, the need to treat the underlying disease that precipitated the stenosis is not Most activities requiring lifting and straining, such as weight changed after a revascularization procedure. 32 Despite the training, have a large static component. In such activities, there is revascularization, some patients are anxious about resuming increased peripheral vascular resistance, with an expected increase physical activity after percutaneous coronary intervention and need in blood pressure but much less of an increase in heart rate or cardiac supervised cardiac reha bilitation to enhance their confidence to output compared with aerobic exercise. Nevertheless, brief periods undertake physical activity and other favorable lifestyle changes. of moderate resistive exercise appear safe and may pose less of a Supervised cardiac rehabilitation also promotes early identification cardiac burden than aerobic exercises of similar effort. In fact, of new signs and symptoms indicating possible restenosis, leading

Despite the expanded use of percutaneous coronary exercises. had undergone percutaneous coronary interventions were 33 Gradual involvement in resistance training may therefore be randomly assigned to a behaviorally oriented intervention or a beneficial and desirable, especially for patients whose jobs or control group. After 12 months, the intervention patients, compared with controls, improved significantly on self-rated measures of smoking, exercise, and diet habits. Patients also lost weight, improved their exercise capacity, and experienced less chest pain during exertion. Although the mechanisms for decreased mortality with exercise have not been fully explained, exercise training improves the lipid profile, ^{37,38} reduces blood pressure, ^{38,39} lowers fasting glucose concentration, 38,40 and reduces body fat and increases lean body mass. 15,37,41

Exercise training has been shown to reduce mortality in patients after myocardial infarction. In one meta-analysis on the combined results of 10 randomized clinical trials that included 4347 patients (control, 2145 patients; rehabilitation, 2202 patients), the pooled odds ratios of 0.76 for all-cause death and of 0.75 for cardiovascular death were significantly lower in the rehabilitation group than in the cardiovascular disease. 30 In addition, patients who have undergone control group, with no significant difference for nonfatal recurrent

In a later meta-analysis of the combined results of 48 clinical trials with 8940 patients, there was a similar 25% reduction in cardiovascular mortality but no significant difference in the rates of nonfatal myocardial infarction and revascularization. 43 This beneficial effect of cardiac rehabilitation on mortality was independent of coronary heart disease diagnosis, type of cardiac rehabilitation, dose of exercise intervention, length of follow-up, trial quality, and trial publication date, which was through March

Patients with heart failure often experience fatigue and dyspnea with exertion. Although the primary pathology of heart failure results from cardiovascular dysfunction, abnormalities in peripheral blood flow, skeletal muscle morphology, metabolism, strength, and endurance all contribute to the heart failure syndrome. ³² Rest was frequently recommended for patients with heart failure in the past, 44 but it is now established that exercise training produces substantial physiological benefits, attenuation of symptoms, and improved quality of life for patients with left ventricular dysfunction and chronic heart failure. 32,45-49

Several mechanisms contribute to improved functional capacity in patients with heart failure who participated in

the blood, and improved neu rohumoral axis. These adaptations 51,52 result in increased oxygen delivery or use in the metabolically more active skeletal muscle, thus delaying reliance on anaerobic or, in some instances, an activity in which they have not participated metabolism. Exercise training has beneficial effects on skeletal for many years. Furthermore, for older persons with coronary heart muscle by improving functional, histological, and biochemical - disease, the clinical manifestations represent the effects of the characteristics and by reducing the activation of the muscle neural disease superimposed on the physiological effects of age, which too afferents known as ergoreceptors.

Adverse events related to exercise training in heart failure in activity. published studies have been few. A limitation in the existing improves quality of life. 44.50

Participants in HF-ACTION were randomized from April 2003 had such a hospitalization despite not undergoing exercise fraining. factors. 23,60,62

Among patients in the exercise group, 759 (65%) experienced a fibrillation or flutter), exercise training reduced the incidence of all- cause mortality in both older men and women. cause mortality or hospitalization by 11% (P = 0.03). Participants in exercise training also achieved significant improvements in cardiopulmonary exercise test parameters and distance in the 6- ROLE OF EXERCISE TRAINING IN PATIENTS WITH minute walk test.

Exercise training also conferred modest but statistically significant improvements in self-reported health status compared - ARTERIAL DISEASE with usual care without training. 50 Improvements occurred early and persisted over time. Based on the safety of exercise, improvements in quality of life and functional work capacity, and modest reductions in clinical events, HF-ACTION supports a prescribed exercise training program for patients with chronic heart failure above and beyond usual medical care.

CARDIAC REHABILITATION IN THE ELDERLY

Increasing evidence has accumulated during the past 3 decades that elderly individuals with coronary heart disease can benefit greatly from exercise training and other aspects of cardiac rehabilitation and secondary prevention programs. 23 This is especially important given that those > 65 years of age represent the largest number of individuals with heart disease. 19 Traditionally, components of secondary prevention programming (exercise; smoking cessation; management of dyslipidemia, hypertension, diabetes, and weight; and interventions directed at return to work and psychosocial issues) are provided by the clinician through the office setting or

exercise training and cardiac rehabilitation. 32 Central hemo dynamic through cardiac rehabilitation programs. Cardiac rehabilitation mechanisms include increases in peak cardiac output, heart rate, and programs are particularly well suited to the provision of secondary stroke volume. Peripheral vascular and metabolic mechanisms prevention services, but unfortunately, many older patients who include improved endothelial vasodilator function, increased would derive benefit from these interventions do not participate cellular oxidative enzyme activity, greater oxygen extraction from because of lack of referral or a variety of societal and other barriers.

> A structured exercise program may be novel to older individuals often lead to decreases in exercise capacity and overall physical

Whereas exertional angina pectoris remains common in this age literature is that most studies have been relatively small and have group, an increased percentage of older patients have atypical not been adequately powered to evaluate mortality and morbidity. manifestations of myocardial ischemia, including dyspnea on To examine the issue of exercise safety and effectiveness in a large exertion and poor functional capacity, often exacerbated by sample of patients with heart failure, Heart Failure: A Controlled comorbidities such as chronic lung disease, peripheral arterial 33 Trial Investigating Outcomes of Exercise Training (HF-ACTION) disease, arthritis, and neuromuscular disorders that may limit was undertaken to determine whether aerobic-type exercise training ambulation. Thus, the absence of exercise-related anginal symptoms reduces all-cause mortality or all-cause hospitalization and in older patients with suspected coronary heart disease may merely reflect the lack of physical activity. 53,54

As a means of increasing physical activity and fitness for older 05 through February 2007 at 82 centers within the United States, persons with heart disease, the prescription of exercise is an essential Canada, and France, and the median follow-up was 30 months. The component of secondary prevention. 20,55 The basis for the exercise no 2331 participants were medically stable patients with heart failure intervention in these patients includes improved functional capacity of and reduced ejection fraction. The interventions were usual care with reduced activity-related abnormal signs or symptoms, plus aerobic exercise training (n = 1172), consisting of 36 supervised including fatigue. 52,56-61 Expected outcomes are similar to those for era sessions followed by home-based training, and usual care alone (n = younger patients, although absolute levels of functional capacity in n 1159). Overall, the performance of exercise training was well the elderly are less, and results may require longer program in tolerated and safe, with only 37 patients in the exercise training participation in this age group. 52,56 As with younger patients, a two group having at least one hospitalization due to an event that multidisciplinary approach to secondary prevention that includes occurred during or within 3 hours of exercise, whereas 22 patients exercise may have a positive impact on other heart disease risk

Whether exercise as a part of secondary prevention is associated primary clinical event compared with 796 (68%) in the usual care with a reduction in morbidity or mortality as in younger patients group. In the primary analysis adjusted for heart failure etiology, has yet to be established. However, studies of older patients with exercise training resulted in a nonsignificant reduction in all-cause and without heart disease have suggested a positive impact of to mortality or hospitalization by an absolute reduction of 4%. physical activity on mortality. 63-65 Findings have suggested that However, after adjustment for key baseline characteristics that are light to moderate activity is associated with a significantly lower risk prognostic for these clinical endpoints (exercise test duration, left of all-cause mortality in persons with established coronary heart 7 ventricular ejection fraction, depression, and history of atrial disease and an inverse association between physical activity and all-

PERIPHERAL

Exercise training plays a critical role as a primary treatment of patients with peripheral arterial disease (PAD), with the goal of improving quality of life and functional capacity. According to the practical guidelines of the American College of Cardiology and the American Heart Association (ACC/AHA) for the management of PAD, supervised exercise training is rated as Class I, Level of Evidence A, for the initial treatment of claudication. 66 The first randomized controlled trial was published in 1966; a marked improvement in walking ability was seen in patients with claudication assigned to daily exercise. 67

546 Several prospective randomized trials have demonstrated that exercise training improves claudication symptoms. 68-70 Studies differ in terms of the magnitude of the reported response to exercise, a finding likely explained by variability both in the exercise intervention itself (duration, frequency, and intensity) and in outcome measures. 71 A meta-analysis of 21 nonrandomized and randomized studies reported a 179% increase in pain-free walking distance and a 122% increase in approximately 50% to 200%. 70

subjects found a significant improvement in 33 maximum PAD patients remains unknown. treadmill walking distances in supervised compared with

exercise training angioplasty group even though the ankle-brachial indices in this area is warranted. improved with angioplasty. 73 Subsequently, the same group program but was given "exercise advice" to walk at home.

A trial from the Netherlands randomized 151 patients with to provide more definitive data on the optimal approach to ischemia. treatment of aortoiliac disease.

improved oxygen extraction by the muscle, better walking between the results at 3 months and at 1, 2, or 3 years. efficiency from a biomechanical standpoint, and enhanced endothelial function. ^{79,80} Calf skeletal muscle may increase its oxidative capacity; improvements in exercise have correlated with changes in carnitine metabolism. 78

In addition, the benefits of exercise may not be fully explained by local effects on the lower extremities, as evidenced by a randomized trial in which 104 patients with PAD were assigned to upper limb aerobic exercise, lower limb aerobic exercise, or no exercise for 24 weeks. 81 Similar improvements in claudication distance and maximum walking distance were observed in both exercise groups but not in the control group, suggesting a systemic benefit of exercise; an increase in exercise pain tolerance was also noted in the exercise groups.

Exercise may attenuate the inflammatory response in the long maximum walking distance. 68 A more recent Cochrane term 71 and has well-established beneficial effects on cardiovascular review of 22 randomized controlled trials for claudication risk factors such as hypertension, diabetes, dyslipidemia, and reported an overall improvement in walking ability of obesity. Further rigorous prospective investigation is needed to ascertain whether this global improvement in vascular health may Overall, patients can expect approximately a doubling of translate into reduced morbidity or mortality in patients with PAD. their pain-free and maximum walking distances with exercise A retrospective cohort study from Japan showed that completion of training, with the best results occurring in supervised programs a 12-week supervised exercise training program reduced both compared with unsupervised home-based training. Another cardiovascular morbidity and mortality in patients with PAD. 82 The Cochrane review of eight small trials including a total of 319 long-term impact of supervised exercise training on outcomes in

Whereas the efficacy of exercise training for the treatment of unsupervised programs, with a difference of approximately 150 symptomatic PAD has been well established, recent evidence now meters. 72 The magnitude and durability of the benefits of home-suggests that exercise training may also play an important role in the based exercise for treatment of claudication, however, remain treatment of asymptomatic PAD. McDermott and coworkers 79 found that supervised treadmill training improved walking and quality of Surprisingly, only a small number of studies have life in patients with PAD with and without claudication. Lower with lower extremity extremity resistance training was also beneficial for improving revascularization for the treatment of claudication. One such functional performance, but not to the same degree as supervised prospective randomized trial compared supervised exercise treadmill training. Given that the large majority of patients with with angioplasty and found that mean maximum walking PAD are either asymptomatic or have atypical symptoms, these distances progressively increased in the supervised exercise findings are particularly important and may herald a significant group at 6, 9, and 12 months but did not increase in the change in the clinical approach to treatment of PAD; further research

In spite of the numerous studies demonstrating the efficacy of reported that after a median follow-up of 70 months, the exercise training, several barriers prevent its wide spread use. functional outcome was the same between the exercise and Perhaps most notably, supervised exercise is not generally a covered angioplasty groups. 74 Among 62 patients randomized to benefit under most health insurance plans including Medicare, angioplasty or medical treatment, 75 there were no significant although PAD rehabilitation has had a Current Procedural group differences at 2 years in treadmill walking distances or Terminology code (CPT 93668) since 2001. Other barriers to quality of life; the medical group was not in a supervised participation include a time commitment and lack of broad availability of PAD rehabilitation centers.

Barriers also exist at the provider level, as many clinicians may claudication to either endovascular revascularization hesitate to refer a high-risk patient with multiple comorbidities to an (angioplasty with conditional stenting) or 24 weeks of twice- exercise program. However, that same patient may in fact derive the weekly 30-minute hospital-based treadmill exercise sessions . greatest benefit from exercise. Because of the systemic nature of ⁷⁶ Endovascular revascularization provided more immediate obstructive atherosclerotic vascular disease and the high prevalence clinical success, but after 6 and 12 months, the treatment of coronary artery disease in this population, patients with PAD in groups were equivalent in terms of functional capacity and many cases may qualify for cardiac rehabilitation from an insurance quality of life scores. The CLEVER (Claudication: Exercise standpoint by meeting one of the other criteria as noted elsewhere Versus Endoluminal Revascularization) trial is a similar in this chapter (eg, recent myocardial infarction, coronary ongoing National Institutes of Health/National Heart, Lung, revascularization, or angina). 29 The decision to enroll a patient with and Blood Institute-funded multicenter randomized clinical PAD in an exercise training program should be individualized and trial comparing supervised exercise with endovascular may be affected by medical comorbidities that limit walking ability, revascularization for the treatment of claudication due to such as pulmonary or degenerative joint disease. Treadmill exercise aortoiliac disease. 77 The results of the CLEVER trial are hoped is not recommended in patients with foot ulcers or critical limb

A meta-analysis found that the best results were seen with The exact mechanism to explain why exercise training walking programs lasting more than 30 minutes per session, at least improves claudication symptoms is not known but is likely three times per week, for at least 6 months. 68 Studies suggest that multifactorial. Initial theories proposed that exercise increases the benefits of the supervised program may extend beyond the collateral vessels through angiogenesis or leads to increased duration of the program. For example, among patients who blood flow, but subsequent data on these hypotheses are attended supervised training twice weekly for 10 weeks, the inconsistent. 71,78 More likely, mechanisms for the benefit of improvements in claudication distance and maximum walking exercise training include metabolic adaptations with distance were sustained at 3 years 83; there was no difference

EXERCISE TESTING FOR PERIPHERAL ARTERIAL DISEASE

Per the ACC/AHA guidelines, standardized treadmill testing should ideally be performed before initiation of exercise training to establish the magnitude of the functional limitation due to claudication and to provide a baseline for measuring the response to therapy. 66 The treadmill test may reveal nonvascular factors Patient contributing to the impaired walking ability (such as degenerative Methods for prescribing exercise for cardiac patients generally do joint disease) and assist in individualizing the exercise prescription. Continuous electrocardiographic monitoring is useful, particularly given the high prevalence of concomitant coronary artery disease.

A graded treadmill test is preferable to a constant-load test because it is more reproducible and shows less variability. 84 For example, the Gardner protocol starts at 2 mph at a 0% degree; the grade is then increased by 2% every 2 minutes. 85 Data including the time at onset of claudication symptoms and the maximum walking time should be recorded. A 6-minute walk test may be used as an alternative to treadmill testing. 66.86

The need for formal cardiac testing before initiation of a PAD exercise program should be individualized. Exercise centers differ in terms of enrollment requirements. At present, a limited though growing number of clinical exercise centers in the United States offer PAD rehabilitation programs. At least at most centers, the patient will need a referral from a physician and a baseline electrocardiogram. Preferably, the referral should also include a recent clinical note and the results of a graded exercise test to establish the cardio vascular response to exercise. Sessions are supervised with monitoring of heart rate and blood pressure; electrocardio graphic telemetry is recommended for at least the first session and may be used throughout each session if necessary. Patients with diabetes require blood glucose monitoring to avoid hypoglycemia as exercise increases insulin sensitivity. In addition, proper footwear and routine self-examination of the feet are important, particularly in patients with diabetic neuropathy.

GUIDELINES FOR EXERCISE PRESCRIPTION INCLUDING THE ROLE OF AEROBIC AND RESISTANCE TRAINING

Healthy Adults

Most epidemiological studies confirm significant risk reduction for those achieving at least moderate-intensity physical activity on most days of the week compared with those who are sedentary. 3 Further increases in fitness produce relatively modest additional muscle fiber hypertrophy (although limited), and increased benefits. Thus, individuals do not need to attain high levels of fitness to accrue substantial health benefits from exercise.

American Heart Association updated their physical activity resistance training significantly increases strength, gait velocity, guidelines for adults. 1 To promote and to maintain health, all balance and coordination, walking endurance, and stairhealthy adults aged 18 to 65 years old need moderate-intensity climbing power. 88-90 aerobic physical activity for a minimum of 30 minutes on 5 days each week or vigorous-intensity aerobic physical activity for a minimum of 20 minutes on 3 days each week. Combinations of moderate- and vigorous-intensity activity can be performed to Aerobic forms of exercise training are designed to increase meet this recommendation. For example, a person can meet the functional capacity and endurance and to increase overall energy recommendation by walking briskly for 30 minutes twice during expenditure while enhancing quality of life. The primary the week and then jogging for 20 minutes on two other days.

Moderate-intensity aerobic activity, which is equivalent to a brisk walk and noticeably accelerates the heart rate, can be

accumulated towards the 30-minute minimum by performing 547 bouts each lasting 10 minutes or more. Vigorous-intensity activity is exemplified by jogging and causes rapid breathing and a substantial increase in heart rate. In addition, every adult should perform activities that maintain or increase muscle strength and endurance a minimum of 2 days each week. Because of the doseresponse relationship between physical activity and health, persons who wish to further improve their personal fitness, reduce their

risk for chronic diseases and disabilities, or prevent unhealthy weight gain may benefit by exceeding the minimum recommended amounts of physical activity.

Exercise Prescription Guidelines for Patients with Cardiovascular Disease Including the Older

not require significant modification for older patients. The exercise prescription should define individual patient guidelines for activity while promoting variety in the exercise regimen. The exercise prescription should encourage the participant to engage in all aspects of physical fitness, including aerobic capacity and muscle endurance, range of motion and flexibility, and muscle strength. Modification of the components of the exercise prescription should be routinely considered, particularly for those persons with comorbidities that limit mobility (eg, musculoskeletal limitations, arthritis, pulmonary disease, and peripheral arterial disease). 33 Increase in energy expenditure and enhancement of functional

independence should be emphasized, as well as participation in activities that increase socialization, which should affect feelings of isolation and depression, not uncommon in this age group. Increasing frequency and duration of activity should supersede increases in intensity, thereby reducing the likelihood of overuse

Recommendations for increasing participation in programs of physical activity should include a broader interpretation of exercise programming, including vocational and recreational activities as well as activities of daily living. Recommendations should also be sensitive to considerations of differences in needs between women and men and ethnic and racial diversity. Because the likelihood of participation is increased with physician referral and support, it is still incumbent upon clinicians to strongly and repeatedly encourage participation in a supervised exercise program.

The use of strength training and flexibility activities for all patients as integral components of the overall exercise prescription should also assist in improving neuromuscular function, muscle strength, and endurance. Such training is essential to improvement of responses to the various physical demands of daily living as well as occupational and recreational activities and improvement of functional independence, selfesteem, and health-related quality of life. In men and women, strength training improves through neuromuscular adaptation, muscle oxidative capacity due to the combination of aerobic exercise and strength development, which is a feature of In 2007, the American College of Sports Medicine and the resistance-type circuit training. 87 Even in the oldest persons,

Aerobic Training

components of the aerobic exercise prescription are

TABLE 33—2 Recommended Components of the Aerobic Exercise Prescription

Frequency

3 to 5 sessions/week initially, progressing to every day when appropriate

Intensity

Heart rate reserve: Define a target heart rate based on peak exercise heart rate, resting heart rate, and a selected percentage based on fitness level (50% to 60% initially, progressing to > 70% as indicated).

Percentage of peak heart rate: Select a target percentage of peak exercise heart rate (60% to 70% initially, progressing to 85% as indicated).

Percentage of peak MET level: Select a target percentage of peak functional capacity in METs (50% to 60% initially, progressing to > 70% as indicated).

Perception of exertion: On the Borg Scale of Perceived Exertion (rating of 6 to 20), patients should attempt to achieve a perceived exertion level of 11 to 16. It is particularly useful for patients who have difficulty measuring heart rate or as an adjunct to heart rate measurement.

Duration

33 5 to 20 minutes initially, depending on intensity of effort and the patient's tolerance, progressing to 30 to 40 minutes as indicated

Modality

Exercise activities should use large muscle groups, that is, legs and arms alone or in combination (eg, outdoor or treadmill walking, cycling, arm ergometry, rowing,

Avoid high-impact activities (eg, jogging), particularly early in the exercise program, to reduce the potential for injury

Exercise may be continuous (few, if any, short rest periods during the exercise session) or discontinuous (periodic rest periods during the exercise session).

Progression

Changes in the exercise prescription should be based on the patient's tolerance to activity. Signs or symptoms of intolerance (eg, angina, significant cardiac arrhythmias, abnormal blood pressure response, unusual shortness of breath, or muscle/orthopedic distress) require evaluation and possible reduction in intensity or duration of exercise.

Conversely, patients may periodically receive increases in exercise intensity or duration of activity as exercise training responses improve (eg, reduced exercise heart rate and systolic blood pressure, as well as perception of effort to standardized exercise workloads)

From the American College of Sports Medicine: Thompson WR, Gordon NF, Pescatello LS: ACSM's guidelines for exercise testing and prescription, ed 8, Philadelphia, 2010, Lippincott Williams & Wilkins.

interrelated and include frequency, intensity, and duration. As an exercise in older cardiac patients.

accomplished by less specific programming, which may include muscle groups. 92 any number of daily vocational and recreational activities Estimates of various levels of energy expenditure are available that initiation of resistance training after a cardiac event are available, can assist both patients and physicians in making decisions about conventional guidelines suggest a restrictive maximal weight limit which activities are appropriate. An example is found in Table 33- (of 10 to 20 pounds) for up to 12 to 16 weeks, particularly in 3, in which estimated metabolic equivalent (MET) levels are unsupervised activities. However, Stewart and colleagues 93 described in multiples of resting oxygen uptake. However, these initiated resistance training as part of a combined resistance and estimates of energy expenditure can be highly disparate from aerobic program as soon as 6 weeks after myocardial infarction, actual values, depending on the individual's level of vigor.

Resistance Training

Aging skeletal muscle responds to progressive overload through resistance training. However, emphasis must be

TABLE 33—3 Examples of Estimated MET Levels for Various Activities			
Specific Activity	Estimated MET Level		
Bicycling for leisure (light to moderate effort)	4-8		

Conditioning exercises (light to moderate effort)	3-7
Home activities (light to moderate effort)	2-6
Lawn/garden activities (light to moderate effort)	2-6
Fishing and hunting	2-6
Jogging/running (light to moderate effort)	7-8
Self-care	2-4
Sexual activity	1-2
Various sports activities	2-10
Walking 2.0 mph	2-3
Walking 3.0 mph	3-4
Walking 4.0 mph	4-5
Swimming for leisure (light to moderate effort)	6-8

From Ainsworth BE, Haskell WL, Whitt MC, et al: Compendium of physical activities: an update of activity codes and MET intensities, MedSci Sports Exerc 32(Suppl):S498, 2000.

placed on providing older persons with adequate time for musculoskeletal adaptation and the development of appropriate training technique, particularly at the initiation of a resistance training program; this practice will reduce the likelihood of muscle overuse, soreness, and injury. 31 This is important in older patients with cardiovascular disease and is also essential for those with hypertension, arthritis, pulmonary disease, or other physically limiting conditions. 91

The initial workload intensity and frequency of training should be modest, providing for the use of proper body mechanics and avoidance of breath-holding and straining during exercise. The initial exercise prescription and subsequent progression of resistance should be undertaken with caution. Alternatives to traditional resistance can be considered , including aquatic resistance exercise and modification of the exercise components, such as variation of exercises, more gradual progression, and increased rest periods within each session.

Table 33-4 describes recommendations for the prescription of example, when the intensity of the activity is limited, as is often the resistance training. In general, the methods and considerations for case with older patients with cardiovascular disease who have prescribing resistance in older adults are not different from those reduced functional capacity or various comorbid conditions, the in younger persons. However, as mentioned previously, frequency or duration of the activity might be increased. Table 33- modifications may be needed to accommodate health conditions 2 provides a general schematic for the prescription of aerobic and other individual limitations. 31 Because the effects of any exercise program including resistance are specific to the muscle Increasing overall levels of physical activity may also be groups being trained, training regimens should involve all major

> Although few data identifying the appropriate timing for demonstrating significantly improved functional capacity and arm and leg strength with no adverse clinical or exercise-related events. As the resistance partici pant progresses in the training program, resistance intensity may be increased to provide for further improvement. In

TABLE 33—4 Prescription of Resistance Training for Older Adults (> 50 years)

Select 6-8 Different Exercises That Involve Major Muscle Groups (Examples)

Upper body: chest press, shoulder press, triceps extension, biceps curl, and pull-down (upper back)

Midsection of the body: lower back extension and abdominal crunch/ curl-up Lower body: quadriceps extension or leg press, leg curls (hamstrings), and calf raise

Exercise Prescription Components

Initially, use single sets of 6-8 varied exercises, 2 days per week.

Each set should include 10-15 repetitions of an exercise at < 40% of 1-repetition maximum (1-RM). §§§§§§§§

Provide 1-2 minutes rest between exercises.

Alternate between upper and lower body work.

Exercise Technique

Perform each exercise through a full range of motion, in a controlled rhythmic manner, at a slow to moderate speed.

Avoid breath-holding and straining (Valsalva maneuver) by exhaling during the contraction or exertion phase of the exercise and inhaling during the relaxation or rest phase of the exercise.

Emphasize proper body mechanics throughout each exercise.

Exercise Progression

As 12-15 repetitions of a given exercise become easily accomplished, consider updating the prescription in the following order:

Different types of exercises.

Increase level of resistance: increasing by 5% of 1-RM, up to 40% 1-RM for arm exercises and up to 60% 1-RM for leg exercises.

Add a second set of exercises without increasing the level of resistance.

Add a third day of resistance training during the week

EXERCISE PRESCRIPTION FOR PERIPHERAL ARTERIAL DISEASE

A standardized program consists of three sessions per week for 12 weeks, although more recent studies have employed a 24-week program. ^{76,79} The participant starts with 5 minutes of warm-up, followed by 50 minutes of intermittent exercise and 5 minutes of cool-down. The treadmill typically starts at 0% grade at 2 mph; the grade is kept constant during the course of each session. A graded pain scale of 1 to 5 is used to rate the claudication symptoms to allow the exercise physiologist to monitor the progress. ⁹⁷

The participant walks on the treadmill until the claudication - discomfort reaches a moderate level (4), preferably within the first 5 minutes on the treadmill. He or she then gets off the treadmill and rests in a chair until the discomfort completely subsides. Treadmill walking then begins again, with repeated bouts of exercise and rest, ideally for a total of 50 minutes.

To maximize the benefit of the exercise training sessions, the grade and speed are increased as walking ability improves. When the patient is able to walk for 5 to 8 minutes at the current speed and grade without stopping, the settings should be changed for the next session. For example, the grade may be increased by 2% per session up to 10%; the speed may then be progressively increased as tolerated. Patients should also be instructed to walk at home for 30 to 60 minutes at least twice per week in addition to the three supervised sessions.

Progress in an exercise program should be objectively measured with subsequent treadmill testing, recording variables such as pain-free walking time or distance, maximum walking time or distance, or the work performed in metabolic equivalents calculated from the speed and grade of the tread mill. Questionnaires such as the Walking Impairment Questionnaire are not typically used in clinical practice but may be employed as time permits to assess functional status. ⁸⁴

Repeated testing with ankle-brachial indices is not generally necessary as the results typically do not change in spite of the expected improvement in functional capacity. ⁷¹ As claudication symptoms improve and exercise tolerance increases, patients should be monitored closely for any signs or symptoms of cardiac ischemia, which may have initially been masked by a limited walking ability.

After the completion of a supervised program, patients should be given an exercise prescription for a home-based maintenance program with the use of claudication symptoms as a guide. Homebased exercise should be similar to the supervised program with intermittent periods of exercise to moderate levels of pain followed by rest until the pain completely subsides.

§§§§§§§1-RM is the greatest amount of resistance (ie, weight) that can be lifted or pushed with a single effort. As an alternative approach, a method of trial and error to determine a level of resistance for which the participant can perform 10 repetitions can be used. The participant can thus work towards comfortably progressing to 15 repetitions before increasing resistance.

older persons, this increase can be achieved by increasing the resistance (or weight) or adding a second set per exercise. Increasing the number of repetitions within a set or decreasing the duration of rest periods between sets or exercises is generally not recommended for this age group. ³¹

Cardiovascular responses to resistance training should be measured, including heart rate, blood pressure, and perception of exertion. Because of the short duration of individual resistance exercises, heart rate response is generally lower than during aerobic exercise. However, blood pressure response can be greater, and thus heart rate alone may not accurately reflect overall cardiovascular response.

In patients who have a history of hypertension, blood pressure sure response to resistance training should be evaluated, especially in those who have known heart disease. However, blood pressure measured immediately after rather than during the actual resistance exercise is likely to underestimate the pressure response. ⁹⁵ Consequently, blood pressure ideally should be assessed during the last couple of repetitions during resistance training, along with a rating of perceived exertion; a rating of 11 to 14 ("fairly light" to "somewhat hard") on the Borg category scale is recommended . ^{31.96} Participants should be frequently reminded of potential adverse signs and symptoms, such as dizziness, excessive shortness of breath, chest discomfort, heart rhythm irregularities, or acute pain in the muscles or joints; resistance training should be immediately discontinued with these occurrences. ^{31.96}

Several tools may improve compliance at home, such as followup phone calls, using a pedometer, joining a gym, setting goals, keeping a walking diary or logbook, finding an exercise "buddy," and putting daily exercise on a "to-do list." Websites such as http://startwalkingnow.org/ sponsored by the American Heart Association may be used to plot walking routes and to log walking times and distances. Throughout the supervised exercise training and beyond, regular medical follow-up should be encouraged to optimize medical management focused on cardiovascular risk reduction.

LONG-TERM MAINTENANCE OF TRAINING EXERCISES

Unfortunately, long-term adherence to formal exercise programs remains a continuing challenge. Because regular exercise training and participation in physical activity are



550 necessary to maintain the physiological benefits described throughout this chapter, many of these benefits are lost in a few weeks when adherence fails. Asking and advising the patient about exercise and physical activity patterns should be a routine part of each visit.

To further emphasize its importance, health care providers may refer to specific advice about activity as an "exercise prescription," underscoring that it is just as important as medicines that might also be prescribed. Both patients and health care providers should adopt the understanding that "exercise is medicine" in terms of its importance to the restoration of health and prevention of disease.

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CHAPTER 34

Psychological Risk Factors and Coronary Artery Disease: Epidemiology, Pathophysiology, and Management

Alan Rozanski

KEY POINTS

- During the last three decades, epidemiologic studies have demonstrated a consistent relationship between psychosocial variables and coronary artery disease.
- Psychosocial risk factors for coronary artery disease include depression; various anxiety syndromes, such as phobias, panic, and post- traumatic stress syndrome; anger/hostility; negative cognitive patterns such as pessimism; and chronic stress, including work stress, marital stress, social isolation, and low socioeconomic status.
- Important new data indicate that positive psychological factors, including positive emotions and having a sense of purpose, improve physiology and increase longevity.
- The pathophysiology governing these relationships stems from chronic activation of the autonomic nervous system and the hypothalamic- pituitaryadrenal axis.
- Both the brain itself, which is remodeled by chronic stress, and the heart are end-target organs of chronic psychosocial
- Psychosocial factors such as depression and chronic stress lead to widespread peripheral effects that promote coronary artery disease, such as inflammation,

- hypercoagulability, metabolic syndrome and diabetes, hypertension, endothelial dysfunction , and enhanced physiological reactivity to environmental stimuli.
- Acute psychological stress can increase the risk for cardiac events among patients with coronary disease through widespread pathophysiological effects.
- Mixed results among sparse large-scale studies have inhibited the adoption of formal guidelines for the management of psychosocial stress in clinical practice.
- New developments within medical psychology hold promise for optimizing behavioral interventions in the future.

Since antiquity, there has been a strong notion the relationship psychological stress and the development of heart disease. However, it was not until the late 1970s that a significant scientific evidence base began to coalesce in this arena. By the early 1960s, the Framingham study had already identified key clinical risk factors for coronary artery disease (CAD), including smoking, hypertension, hyperlipidemia, diabetes, and a family history of premature CAD. At that time and into the following decade, interest in potential psychosocial risk factors for CAD was mainly dominated by a construct proposed by Friedman and Rosenman. 1 They conceived of a coronarypersonality type Α characterized as hard driving, time impatient, and prone to hostility. Initial research linked type A personality to a higher risk of cardiac disease compared with type B. Whereas this concept proved popular and became part of the cultural vernacular for many years, the construct was abandoned because of the lack of sufficient confirmatory evidence during prospective

Subsequent research, however, began to define strong links between various psychosocial risk factors and CAD. For instance, in 1979, the landmark Alameda County study was published, noting a stepwise gradient between the size of individuals' social network and all-cause mortality. ² New and more sophisticated epidemiologic studies, which for the first time corrected for con current risk factors such as smoking and involving increasing, more representative

sample sizes, began to demonstrate a similar stepwise gradient between the magnitude of depressive symptoms and adverse cardiac outcomes. ³ At the same time, important animal work began to define pathophysio logical links between chronic stress and atherosclerosis. 4 With respect to acute stress, newly available imaging techniques were able to show for the first time a relationship between acute stress and myocardial ischemic perfusion wall motion abnormalities, and coronary vasoconstriction. ³ Since then, a markedly vast literature has developed to link a large variety of psychosocial risk factors to CAD.

This chapter reviews the current state of the epidemiological link between psychosocial risk factors and CAD. Both negative factors that cause CAD and more recently defined positive factors that help buffer against the development of CAD are reviewed. The pathophysiological basis for these links is broadly examined, and treatment -considerations that are relevant to car -diologists are considered.

THE EPIDEMIOLOGY PSYCHOSOCIAL LINKING RISK FACTORS TO CARDIOVASCULAR DISEASES

During the last few decades, a number of negative psychosocial factors have been linked to the development of cardiovascular disease (CVD). More recently, various positive -psychological factors, such as the

BOX 34-1 Psychosocial Factors That Have Been Linked to the Progression or Prevention of Cardiovascular Disease

Negative Thought Patterns and Emotions

Depressive syndromes

Mild to moderate depressive symptoms

Major depression

Hopelessness

Anxiety syndromes

Generalized anxiety disorder

Phobic anxiety

Panic disorder

Post-traumatic stress disorder

Hostility and anger

Worry Pessimism

Chronic Stress

Work stress

Marital stress and dissatisfaction

Social isolation and lack of social support Low socioeconomic status

Foreign caregiver

Adverse childhood experiences

Perceived injustice

Positive Psychological Factors

Positive emotions Optimism

Social support

Sense of purpose

presence of positive emotions and having a sense of purpose, have been demonstrated to protect against CVD and to promote longevity. The psychological factors that have been shown to be epidemiologically linked to CVD are listed in Box 34-1. These factors can be divided into three broad categories: (1) chronic negative thought patterns and emotions; (2) chronic stress; and (3) positive psychosocial factors that promote health and buffer against CAD. Each of these categories is reviewed.

Thought Patterns and Emotions

Thought patterns and emotions are logically linked because they are bidirectionally related. Thoughts commonly generate concomitant emotional responses, but our moods and emotional states can also affect the quality of our thinking. Depression, anxiety, and anger/hostility have been most commonly studied; but in recent years, other thought patterns, such as pessimism/optimism, have also emerged as an important area of study.

Depression

Depression has been particularly studied as a risk factor for CAD, for a variety of clinical reasons. First, depressive symptoms are common in society; major depression, alone, affects approximately 5% of the US population at any time, with an increased frequency in cardiac cohorts. Second, depression is painful and debilitating, leading to loss of productivity and high economic costs. ⁵ Third, various effective quantifiable tools exist to measure depressive symptoms, and epidemiological studies that use these tools have consistently demonstrated that depression is associated with a heightened frequency of atherosclerotic heart disease and adverse clinical events.

Validated standardized scales, such as the Beck Depression Inventory and the Center for Epidemiologic Studies

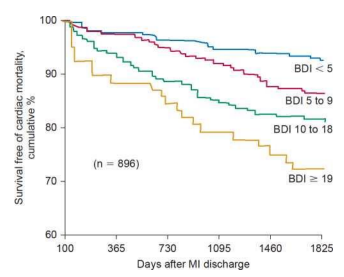


FIGURE 34-1 Grouping of post-myocardial infarction (MI) patients according to their Beck Depression Inventory (BDI) scores, ranging from those with no depressive symptoms (BDI < 5) to those with moderately severe depression (BDI > 19). A gradient relationship was observed for frequency of death according to the magnitude of depressive symptoms. Notably, increased events occurred even among patients with mild depressive symptoms (BDI scores of 5 to 9)

Depression Scale (CES-D), allows depressive symptoms to be characterized along a full spectrum, ranging from mild to severe depressive symptoms. At the severe end of this spectrum is major depression, which is a specific clinical psychiatric disorder that is established by formal diagnostic interview according to the following *Diagnostic and Statisti cal Manual of Mental Disorders*, fourth edition (DSM-IV) criteria: the presence of severely depressed mood and/or inability to take pleasure in all or most things that were previously considered enjoyable, lasting > 2 weeks and accompanied by functional impairment and somatic complaints, such as fatigue or loss of energy nearly every day, insomnia or hypersomnia, change in appetite, diminished ability to concentrate, feelings of worthlessness or inappropriate guilt, and recurrent thoughts of death or suicidal ideation.

Studies have consistently demonstrated that the prevalence of major depression is increased by at least threefold in cardiac populations, occurring in at least 15% to 20% of patients who have post-myocardial infarction status, unstable angina, angioplasty, bypass surgery, or valve surgery or who have congestive heart failure. ³ In addition, at least another 15% of cardiac patients can be expected to manifest minor to moderate depressive symptoms that do not meet the criteria for major depression according to DSM-IV criteria. This high prevalence of depressive symptoms in cardiac populations is believed to be due to a bidirectional relationship between depression and CAD.

This high frequency of depressive symptoms in cardiac patients is of particular concern given repeated evidence of a strong gradient relationship between the frequency of depressive symptoms and the occurrence of adverse cardiac events both among community cohorts who are observed prospectively and among cardiac disease cohorts. Evidence indicates that even mild depressive symptoms are associated with an increased frequency of cardiac events compared with patients who have no depressive symptoms ⁶ (Fig. 34-1).

Various meta-analyses have been performed, and each has found a prognostic association between depression and adverse outcomes. ⁷⁻¹¹ For example, meta-analyses of community-based cohorts conducted by Rugulies ⁷ and by Wulsin and Singal ⁹ found depression to be associated with a relative risk ratio of 1.64 for the development of CAD. Simi larly, Van Melle and coworkers ¹⁰ and Barth and colleagues ¹¹



Study, Year	No. of Subjects	Follow-up	Scale	Endpoints	Adjusted Risk Ratios
McCarron et al,13 2003	9,239	20 years	Physician impression	ACM	1.36 (1.07-1.72)
Eaker et al,14 2005	3,682	10 years	Framingham anxiety scale	ACM, male ACM, female	1.22 (1.08-1.38) 1.27 (1.05-1.55)
Ringback Weitoft and Rosen, 2005 (3 different cohorts)	10,733 10,035 9,100	5 years	Hospital ICD codes	ACM ACM ACM	1.71 (1.1-2.5) 2.3 (1.6-3.3) 1.8 (1.1-1.3)
Ostir and Goodwin, 16 2006	506	5 years	^{Zun} g	ACM	1.52 (1.02-2.28)
Mykletun et al, 17 2007	61,349	4.4 years	HADS	ACM CD	0.90 (0.83-0.98) 0.89 (0.67-3.38)
Fan et al,18 2008	129,499	N/A	History of anxiety on PHQ-8	Self-reported CVD	1.46 (1.37-1.54)
Shen et al, 19 2008	735	12.4 years	MMPI anxiety subscales	MI	1.43 (1.17-1.75)
Phillips et al,2009	4,256	15 years	DSM-III (interview)	ACM CD	1.80 (1.16-2.80) 1.84 (0.98-3.45)

ACM, all-cause mortality; CD, cardiac death; CVD, cardiovascular disease; HADS, Hospital Anxiety and Depression Scale; MI, myocardial infarction; MMPI, Minnesota Multiphasic Personality Inventory; PHQ, Patient Health Questionnaire.

found depression to be associated with at least a twofold increase in event risk for cardiac populations among meta analyzes involving 22 studies and 20 studies, respectively.

Anxiety Syndromes

Like depression, feelings of anxiety vary across a wide spectrum , but anxiety disorders are more common than depression . For instance, within the National Comorbidity Survey, the 12-month prevalence of diagnosed anxiety disorders within a representative national survey was approximately 20%. ¹² Transient experience of anxiety is a universal phe nomenon that is often an adaptive warning of threat or danger. When anxiety is chronic or cannot be controlled, however, it becomes maladaptive. Many studies have now assessed the epidemiological significance of both self-reported anxious symptoms and various pathological forms of anxiety, as characterized in the DSM-IV.

In early studies, conflicting results were obtained regarding anxiety as a CVD risk factor, but in recent years, increasing studies have clarified the status of anxiety syndromes relative to cardiac risk. Table 34-1 lists the outcomes associated with measurements of anxiety in community samples for studies reported since 2003. 13-20 In general, these studies found anxiety to be associated with significant risk ratios for adverse clinical events, although in some studies, anxiety was not a significant predictor after full adjustment for covariates of risk. 13,20 This sensitivity to covariate adjustment may help explain some of the conflicting data concerning anxiety among epidemiological studies.

There have also been many studies concerning anxiety within cardiac populations, but most of these have involved small sample sizes. Among three studies with cardiac populations > 500 individuals, each demonstrated an increased risk for adverse clinical events in the presence of anxiety. 21-23 In an interesting study in this arena, Frasure-Smith and Lesperance 23 compared self-reported anxiety symptoms versus the identification of generalized anxiety disorder (GAD) according to use of the Structured Clinical Interview for DSM-IV. The diagnostic criteria for GAD include excessive anxiety or worry for more days than not for > 6 months, with difficulty in controlling the worry and symptoms of functional impairment and distress (eg, fatigue, insomnia). The study assessed 804 patients with stable CAD observed for a composite index of major adverse cardiac events. Whereas both self-reports of anxiety and GAD were associated with increased events before covariate adjustment, only GAD remained a significant predictor of risk after covariate adjustment. These results are indicative of a gradient relationship, whereby more patho logical anxiety exerts greater pathophysiological effects.

In addition to GAD, most of the other DSM-IV anxiety syndromes have been assessed relative to their clinical sequelae. Among the anxiety disorders that will be encountered by cardiologists is panic disorder, because the symptoms associated with panic attacks, including chest pain, palpitations, and dyspnea, can lead to frequent emergency department presentations. Panic disorder involves the presence of recurrent and unexpected panic attacks, with the occurrence of anticipatory anxiety or worry and a significant change in behavior related to the attacks. Among studies concerning panic attacks, Smoller and coworkers ²⁴ observed 3369 women for 5.3 years and found panic attacks to be associated with a substantial hazard ratio for the development of CVD and stroke as well as all-cause mortality. Panic disorder was also found to be associated with an increased frequency of acute myocardial infarction in the follow-up of 9641 patients with panic disorder who were matched to 28,923 controls, 25 and similar findings were noted in other large studies. ^{26,27}

The most common of the DSM-IV forms of anxiety disorders is the presence of phobic disorders. The 12-month prevalence of simple phobias was approximately 9% among the individuals within the National Comorbidity Survey, and social phobias were found within approximately 8%. ²⁰ The DSM-IV criteria for phobia include the presence of excessive and persistent fear that is cued by the presence or anticipation of exposure to a specific object or situation, with the exposure usually invoking an immediate anxiety response. Notably, phobic anxiety, as assessed by the Crown-Crisp Index, has also been linked to cardiac death in large cohorts, including the follow-up of 33,999 men from the Health Professionals Follow-up Study ²⁸ and 72,359 women from the Nurses' Health Study. ²⁹

Another anxiety disorder that has recently been strongly linked to cardiovascular sequelae is post-traumatic stress disorder (PTSD). PTSD is considered present if, after the exposure of an inciting traumatic event, subjects report reexperiencing the traumatic event, hyperarousal, and avoidance of traumatic reminders and emotional numbing. In

a prospective study of men who had served in the military, a stepwise relationship was noted between increasing symp toms of PTSD and both cardiac death and nonfatal myocar dial infarction, ³⁰ and a second study found a relationship between PTSD and incident CAD among a community cohort of 1059 women who were observed for 14 years. ³¹ In a third study, Boscarino ³² evaluated 4328 men who had served in the Vietnam War. The presence of PTSD was associated with a more than twofold increase in the frequency of subsequent cardiac mortality, with the effect being independent of the presence of depression. PTSD patients are more prone to depression, but evidence indicates that pathophysiological effects from PTSD, such as hypertension, may occur independently of depressive symptoms. ³³

Anger and Hostility

Anger and hostility are often grouped together in studies of psychological risk factors because of overlapping characteristics . Hostility refers to an entrenched cognitive trait of easily precipitated resentment, cynicism, or suspicious negative thoughts about others. This cognitive style leads to frequent expressions of anger and negative social exchanges. Anger is an acute negative emotion, but individuals may by temperament or due to life experience be predisposed to a cognitive style of angry thoughts that may result in either expressed or suppressed feelings of anger. Hostile individuals are frequently angry, but anger can be experienced without hostility.

Unlike for depression or anxiety, no psychiatric system of classification has been developed for patients manifesting syndromal hostility or anger. Also, research in this arena has been characterized by the use of widely varying scales, which may have contributed to disparate findings over the years. Moreover, whereas anxiety and depression can be assessed both by self-report and by independent structured interview, the use of a standard structured interview approach has not been commonly applied for the study of anger and hostility. This represents a potential limitation in this arena because lack of self-awareness or self-denial may potentially limit the accuracy of self-reports concerning anger and hostility. Nevertheless, despite such limitations, an increasing literature regarding anger and hostility has emerged.

Chida and Steptoe ³⁴ recently conducted a meta-analysis of published prospective cohort studies concerning anger and hostility. Among 25 studies involving community samples, the hazard ratio for CVD events in initially healthy populations in association with anger and hostility was 1.19 (95% CI, 1.05-1.35). Similarly, the hazard ratio was increased to 1.24 (95% CI, 1.08-1.42) among 19 studies involving patients with known CVD. Further support for the link between hostility or anger and CVD comes from a number of studies that have found these psychological variables to be associated with greater presence or progression of objectively measured atherosclerosis. ³

Pessimism Versus Optimism

The tendency that people have towards thinking in pessimistic versus optimistic patterns has been closely linked to health outcomes. Optimism and pessimism have most commonly been measured in medical research in two ways. One characterization, developed by Seligman and colleagues, defines optimism versus pessimism in terms of the way individuals attribute the *causes* to life events. ³⁵ Optimists tend to see negative events as temporary and positive events as more permanent; they tend to attribute specific causes to negative events while viewing positive events in more global terms; and they tend to attribute external causes to negative events rather than employing self-condemnation. Pessimists have the opposite "explanatory style." Another characterization of optimism versus pessimism, as developed by Scheier and



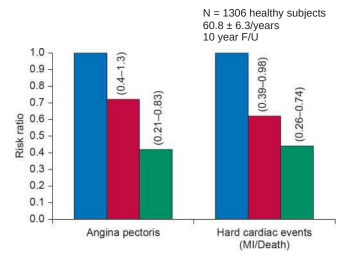


FIGURE 34-2 Occurrence of incident angina and hard cardiac events among 1306 healthy subjects in the Normative Aging Study, followed up for 10 years.³⁷ A gradient relationship was noted for outcomes for subjects classified as having a pessimistic, neutral, or optimistic explanatory speaking style.

Carver, ³⁶ defines optimism versus pessimism in terms of -individuals' disposition toward expecting positive or negative future outcomes. This disposition toward optimism or -pessimism is seen as influencing people's behaviors, how they pursue goals, and their resilience to life stress.

Both approaches towards measuring pessimism and optimism have been linked to adverse clinical events. A 10-year follow-up of 1306 men from the Normative Aging Study found that those with a pessimistic explanatory speaking style had the highest frequency and those with an optimistic speaking style had the lowest frequency of incident angina and coronary events (myocardial infarction or cardiac death) during follow-up, whereas those who had a neutral speaking style had an event rate that was intermediate (Fig. 34-2). ³⁷

In a study of 7216 subjects who were observed during four decades, those who tested pessimistic on a personality inventory showed a significant linear trend toward greater all-cause mortality with increasing pessimism. ³⁸ Among elderly subjects observed for approximately 9 years, those subjects who tested high for optimism, compared with high scores for pessimism, had a substantially reduced hazard ratio for all-cause mortality (0.55; 95% CI, 0.42-0.74) and cardiac mortality (0.23; 95% CI, 0.10-0.55). ³⁹ In another elderly cohort, the degree of measured dispositional optimism was found to be inversely associated with the frequency of cardiovascular death in a cohort of 545 men observed for 15 years. ⁴⁰

Most recently, Tindle and associates ⁴¹ assessed the relationship between optimism and CVD in the largest cohort yet studied, 97,253 women from the Women's Health Initiative. Optimistic women, compared with pessimistic women, again manifested a reduced adjusted risk ratio for cardiac mortality (0.70; 95% CI, 0.55-0.90) in this study. Together, these studies provide consistent evidence for a strong relationship between pessimism and optimism and adverse clinical events.

Worry, Rumination, and Other Negative Thought Patterns

A variety of other negative thought patterns, which may relate to anxiety or depression or serve as precursors to these psychological states, have received attention in the cardiovascular literature but not with a sufficient breadth of epidemiological study to date. One of these cognitive states is chronic worry, which represents a cognitive component or precursor of



Rumination is another negative cognitive state involving the tendency to repetitively think about negative events. Whereas epidemiological study is lacking, ruminators have been shown to have heightened heart and blood pressure reactivity to acute stress and delayed recovery of these responses. 43 Perfectionism also represents a negative cognitive state. Perfectionism is characterized as the tendency to set excessively high standards for performance, accompanied by a highly critical style of selfexamination. Preliminarily, perfectionism has been linked both to mortality in one study 44 and to excessive cortisol secretion during psychological stress in a second study. 45 The cognitive tendency to be for giving or unforgiving is another cognitive pattern that was found to influence the degree of cardiovascular reactivity to stress in one study. 46 Clearly, much more study is needed in this arena, but combined, these studies are suggestive of a broad link between chronic negative cognitive states and cardiovascular sequelae.

Chronic Stress

The study of chronic stress is unique within the context of psychological risk factors for CVD because chronic stress can be readily studied in controlled animal models of stress, complementing the epidemiological study in humans. A set of investigations performed in cynomolgus monkeys (Macaca fascicularis) has been particularly insightful for elucidating the relationship between chronic stress and atherosclerosis. 4.47-49

Cynomolgus monkeys are an apt model for correlative study because they develop coronary atherosclerosis when under stress or when fed fatty diets, with similarities to humans in terms of coronary pathology and pathophysiology. Cynomolgus monkeys also have definable and quantifiable characteristics, and their social environment can easily be modified to create conditions of chronic stress. One approach to inducing chronic stress in these monkeys is to take advantage of the fact that they form well-defined social status hierarchies, with the most dominant monkeys reliably defeating more subordinate monkeys during competitive interactions. Once dominance rank is established, these monkeys form stable hierarchical societies. However, if monkeys are constantly placed into new social groups, a chronically stressful environment is created as the monkeys continually reinitiate their attempt to establish dominance within their new social groups. When male monkeys within stable versus unstable groups were kept on a lowcholesterol diet, the dominant male monkeys within the unstable social groups developed endothelial injury. 4 When both the stable and unstable social groups of monkeys were fed a highcholesterol diet, both groups developed atherosclerosis, but the magnitude of the disease was greater in the dominant male monkeys that were in the unstable environments. 47

Chronic stress has been well studied within the epidemiologic literature. The most commonly studied chronic stress in this regard has been work stress. Estimates as to the frequency of chronic work stress vary widely, but it is a common stressor. The leading models of work stress are the model of job strain as developed by Karasek and associates 50 and the effort-reward imbalance model as developed by Siegrist 51 (Fig. 34-3). The job stranger model posits that individuals experience job stranger when experiencing work that is highly demanding but associated with low job latitude. Lack of control, however, appears to be a more toxic factor than high job demand. 52 In the effort-reward imbalance model, stress

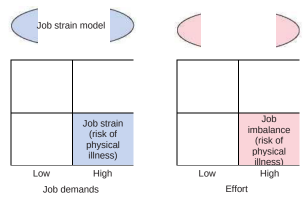


FIGURE 34-3 Two common conceptual models of job stress. The foreign job model (left) is based on assessment of the amount of job demand and decision latitude at work. The presence of high demand but low decision latitude is characteristic of job stress. The effort-reward imbalance model (right) is based on assessment of job demand versus "reward" at work, whether financial or in terms of nonfinancial factors such as recognition, advancement, and prestige at work. High effort with low reward is characteristic of job imbalance. (From RozanskiA, Blumenthal JA, Davidson KW, et al: The epidemiology, pathophysiology, and management of psychosocial risk factors in cardiac practice: the emerging field of behavioral cardiology. J Am Coll Cardiol 45:637, 2005.)

occurs when high demand is associated with low "reward," either in terms of financial or professional reward (eg, opportunities for promotion) or in terms of psychological reward (eg, sense of security or self-esteem). Notably, both models have been linked to adverse CAD outcomes, as summarized in a review of 13 prospective cohort studies. 53 Compared with each other, the job strain and effort-reward imbalance models were comparable predictors for adverse outcome. 54

Other studies have also linked work stress to the presence of accelerated atherosclerosis during serial carotid ultrasound studies. 55 The effects of work stress tend to be accentuated in individuals with lower socioeconomic status and those with lower social support. However, work stress may also affect white-collar and higher socioeconomic status workers with seemingly more autonomous jobs as high work demand or low latitude commonly occurs because of internal psychody namic factors, such as the need for recognition or perfectionistic tendencies.

Although marital stress or dissatisfaction is common in society, for years it was less studied than work stress relative to its association with CVD. Many studies over the years have studied clinical outcomes relative to marital status (eg, remaining married, single, divorced, or widowed), but only recently have investigators begun to focus on outcomes as a function of marital quality. Among community cohorts, the quality of marital communications and marital conflict were found to strongly influence the frequency of adverse cardiac outcomes during a 10year follow-up of 3682 subjects in the Framingham Offspring Study. ⁵⁶ For example, women who self-silenced themselves during marital conflicts were noted to have a fourfold increase in mortality during follow-up. Similarly, an increased frequency of cardiac events was noted among those reporting more negative marital interactions during a 12.2-year follow-up of 9011 British civil servants. 57 Among patients with CAD, Orth-Gomer and coworkers 58 found marital stress to be associated with a 2.9-fold increased risk of recurrent events during a 5-year follow-up of women who were status post-myocardial infarction, and Coyne and associates 59 found that marital quality influences mortality rates among patients with congestive heart failure. Marital quality has also been found to influence the rate of progression of atherosclerosis as measured during both serial carotid ultrasound and coronary artery calcium scanning. 60

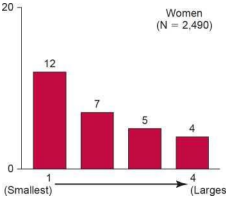


FIGURE 34-4 Mortality in the Alameda County study² according to the size of subjects' reported social network for men and for women, ranging from smallest to largest network size. An inverse gradient was observed in both men and women.

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Data garnered during three decades provide strong epidemiological proof for the health damaging of poor social support and social isolation. Social support in epidemiological studies is commonly divided into the amount of tangible or instrumental support (eg, the size of one's social network) and the amount of emotional social support. Inadequacy in both types of support is linked to a heightened incidence of CVD and reduced longevity, both among community-based cohorts and among patients with known CAD. ³ Within these studies, an inverse gradient has consistently been noted between the amount of social support and the development of CAD or cardiac events ³ (Fig. 34-4).

Two other common stressors are low socioeconomic status and caregiver stress. Low socioeconomic status is associated with such psychosocial factors as higher anxiety resulting from hardship and environmental threat, poorer social support, demoralized feelings, and lower self-esteem. There is strong evidence to suggest that the impact of socioeconomic status on outcomes is due not only to material disadvantage and poorer health habits but also to psychological and stress-related mechanisms. 61-63 By contrast, whereas foreign donor care is commonplace in today's society, it has been much less studied, especially in terms of cardiac outcomes. In the Nurses' Health Study, caring for > 9 hours per week was found to be associated with significantly increased risk of fatal CAD or nonfatal myocardial infarction. ⁶⁴ More study in this arena is required. Recently, other forms of chronic stress have begun to receive attention as potential risk factors for CVD, including the longlasting influence of adverse childhood experiences 65,66 and the experience of organizational injustice. 67

Positive Psychological Factors

Until recently, most investigation regarding psychosocial factors and CVD has focused on just the negative relationship ships associated with factors such as depression and anxiety. However, two psychosocial factors have been persistently associated with a full spectrum of both negative and positive effects: optimism and social support, as reviewed before. More recently, new research has focused on the buffering and health-promoting effects of other positive factors, including positive emotions and the satisfaction of basic psychological needs that can induce a state of relative flourishing. ⁶⁸⁻⁷³ Positive emotions in the context of this research have been defined broadly to include both emotions such as happiness and states of being that reflect a positive engagement with the environment, such as the presence of curiosity and interest. ⁶⁸

In an important work, Fredrickson 68 developed the broadenand-build model concerning emotions. This work has demonstrated an important directional relationship between the quality of emotions and cognitive functions, such as a broader scope of attention, more flexibility, and better problem solving and creativity skills. In positive emotional states, individuals are seen as more likely to be friendly and optimistic. Thus, positive emotion is seen as increasing individuals ' personal resources and providing them with increased resilience to cope with stress. A recent meta-analysis has assessed the relationship between positive well-being and longevity among 35 studies involving initially healthy populations and 35 other studies involving patients. 74 In both cohorts, the presence of positive well-being was associated with reduced mortality. To date, however, there has been little study regarding the effect of positive emotions on cardiac outcomes. Certain attitudinal states have also recently received attention as to their impact on overall well-being. These include the practice of gratitude 75 and social altruism. 76 These parameters, however, have not yet been well assessed in terms of cardiac epidemiology.

In addition to positive emotions, various theorists have postulated that individuals have psychological needs that must be met for optimal satisfaction and that the absence of these needs leads to psychological tension. ^{69,73} Whereas these theorists differ in what constitutes these needs, there is general agreement among virtually all theorists that social connection is a basic psychological need. This may help explain why measured levels of social support have consistently been a very strong disease predictor. In addition, some theorists also posit that individuals have a basic need for meaning or sense of purpose, ^{69,72} which when unmet leads to chronic tension.

Recent data support this assertion. For example, in a study of 1238 elderly individuals, purpose in life was assessed according to a 10-item scale, and the subjects were then observed for a mean of 2.7 years. ⁷⁷ Those who were identified as having a higher level of purpose in life had a substantially adjusted reduced risk of mortality (0.60; 95% CI, 0.42-0.87) (Fig. 34-5). Similarly, in the MacArthur Study of Successful Aging, older adults in the age range of 70 to 79 years who reported being more useful to friends and family during a baseline interview had reduced mortality and disability during a 7-year follow-up compared with those who reported never or rarely feeling useful. ⁷⁸ Similar findings were also noted in a third study involving a 6-year follow-up of elderly subjects in Japan. ⁷⁹ More study is needed to assess the epidemiological effects of having a sense of purpose in younger subjects and its specific relationship to cardiovascular disease.

FIGURE 34-5 Cumulative hazard among 1238 older community-dwelling individuals observed for 5 years, according to the presence of a low versus high sense of purpose. The hazard rate for mortality in persons with high scores for purpose in life was about 57% of that for persons with low scores. (From Boyle PA, Barnes LL, Buchman AS, et al: Purpose in life is associated with mortality among community-dwelling older persons. Psychosom Med 71:575, 2009.)

Positive emotions

Healthy cognition Satisfaction of basic psychological needs



The influence of positive and negative psychological factors as well as the influence of biological and behavioral factors has been postulated to be contained within a composite variable, one's sense of emotional vitality, 80 as conceptualized in Figure 34-6. The emotional aspect of vitality is augmented by positive emotions, positive thought patterns like optimism, and the satisfaction of basic psychological needs like social support and a sense of purpose. By contrast, chronic negative emotions, negative thought patterns like rumination and worry, dissatisfaction of basic psychological needs, and chronic stress are all seen as depleting vitality.

Support for this concept comes from a study by Kubzansky and Thurston, ⁸¹ who assessed the link between the emotional aspects of vitality and incident CAD among 6025 subjects in the National Health and Nutrition Examination Survey (NHANES) I. During a mean follow-up of 15 years, those who manifested higher levels of emotional vitality manifested less CAD.



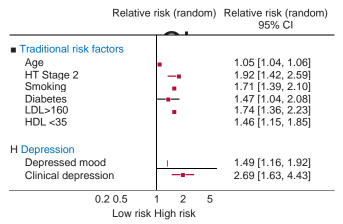


FIGURE 34-7 Comparison of the risk ratios for traditional Framingham risk factors in men versus the risk ratios for depressive symptoms and clinical depression as reported by Rugulies. ⁷The risk ratios for traditional risk factors are for death due to cardiac causes (death, myocardial infarction, coronary artery insufficiency, development of angina). For depressive symptoms and clinical depression, the risk ratios are for death due to cardiac disease and myocardial infarction. Cl, confidence interval; HT, hypertension; HDL, high-density lipoprotein; LDL, low-density lipoprotein. (From Rozanski A, Blumenthal JA, Davidson KW, et al: The epidemiology, pathophysiology, and management of psychosocial risk factors in cardiac practice: the emerging field of behavioral cardiology. J Am Coll Cardiol 45:637, 2005.)

In summary, epidemiological studies have identified an number of psychosocial risk factors I I Energy associated with CAD. Although these risk factors have been studied isolation, verv frequently they cluster together. For example, the Regulation frequency of depression

has been found to be increased *threefold among* individuals experiencing high job stress. ⁸² The risk for CAD and cardiac events associated with many psychological risk factors for CAD generally demonstrates a gradual relationship and a level of risk that is comparable to conventionally studied CAD risk factors (Fig. 34-7).

A unique study that compared psychosocial risk with other CAD risk factors was the INTERHEART case-control study. ⁸³ This study examined a variety of CAD risk factors among an international population of 12,461 acute post myocardial infarction patients, matched to 14,637 control subjects. A simple psychological index in this study was comparable to other CAD risk factors in terms of myocardial infarction risk (Fig. 34-8). This psychosocial index remained a robust predictor of myocardial infarction independent of geographic or ethnic context. ⁸⁴

Of note, within the epidemiological study, the risk associated with psychological risk factors is adjusted for other CAD risk factors. However, because behavioral and metabolic CAD risk factors tend to aggregate disproportionately among individuals with psychosocial stress, in causative fashion, the true cardiovascular risk posed by psychosocial risk factors may be even greater than that reported within the literature.

PATHOPHYSIOLOGY

For many years, the pathophysiological mechanisms responsible for the morbidity and mortality associated with

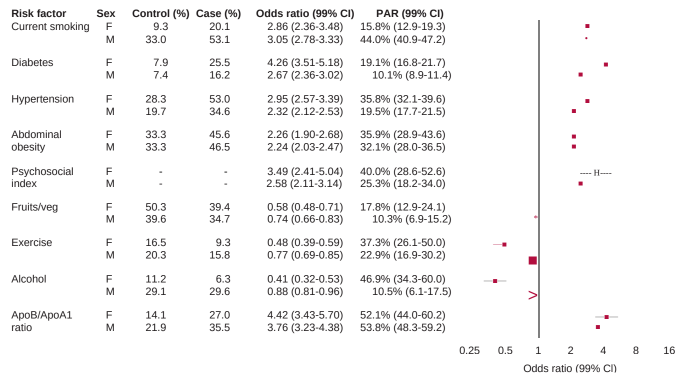


FIGURE 34-8 The risk of acute myocardial infarction results from the international INTERHEART case-control study for each of nine CAD risk factors were evaluated, adjusted for age, gender, and geographic location. The prevalence of each CAD risk factor is presented for controls and cases in the third and fourth columns. The prevalence rates for the psychosocial index were not calculated as it was derived from a statistical odds ratio. PAR, population attributable risk. (From Yusuf S, Hawken S, Ounpuu S, et al: Effect of potentially modifiable risk factors associated with myocardial infarction in 52 countries (the INTERHEART study): case-control study. Lancet 364:937, 2004.)

psychosocial risk factors for CVD were obscure. However, this is no longer the case. Both animal studies and sophisticated research in humans have elucidated a complex patho physiology that is unleashed by negative psychosocial factors, resulting in widespread systemic effects. The pathophysiol ogy associated with depression and chronic stress is summarized in Figure 34-9. These two psychosocial conditions have been most studied relative to pathophysiology and are thus reviewed here. However, there is also ample research into the pathophysiologic effects associated with other common psychosocial risk factors for CVD, such as poor social support, anxiety, and anger/hostility, as well as increasing study of the beneficial effects associated with positive factors such as positive emotions and optimism that are not reviewed in this chapter but are well documented in the literature.

Effects of Chronic Stress

The brain serves as a constant sentinel for all stimuli that are perceived as physically or psychologically threatening in our environment. Perceived threat induces an acute stress response that involves activation of both the autonomic nervous system and the hypothalamic-pituitary-adrenal (HPA) axis. ⁸⁵ When perceived or real threat remains persistent, a chronic stress response ensues, involving profound dysregulation of both systems.

Under conditions of acute stress, the secretion of cortisol serves as a negative feedback system, helping to terminate the physiological response to acute stress. By contrast, chronic stress results in the turn-off of this negative feedback loop, with resulting high cortisol secretion. ⁸⁶ In addition, there is ele vated sympathetic nervous system stimulation during chronic stress. The chronic stimulation of the HPA and sympathetic nervous systems within the brain during chronic stress leads to the cascade of peripheral pathophysiological effects that are outlined in Figure 34-9, including increased inflammation, ⁸⁷

hypercoagulability, 88 metabolic syndrome, 89 central obesity, 90 and hypertension. 91,92

In addition, an important and consistent effect of chronic stress is that it leads to impaired and exaggerated physiological reactivity, as characterized by increased heart rate and blood pressure responses to physiological stimuli. A variety of clinical data indicate that such enhanced physiological reactivity leads to accelerated atherosclerosis. For example, in an experimental study, cynomolgus monkeys exhibiting higher heart rate responses to a threatening experi mental stimulus demonstrated atherosclerotic lesions in coronary and carotid arteries that were twice as large as those of less reactive monkeys. 49 Pretreatment with beta blockade abolished the excess atherosclerosis that is observed in dominant males housed in unstable social environments, indicating that sympathetic activation is an important mediator of this stress-induced atherosclerosis. 48 In parallel to this animal work, individuals with greater physiological reactivity have been found to manifest accelerated atherosclerosis during serial carotid ultrasound study. 3

An area of recent interest is an apparent relationship between chronic stress and premature aging, as assessed by the effect of stress on the length of leukocyte telomeres. Telomeres have repetitive DNA sequences complexed with proteins that cap and protect chromosome ends. They decline in length with age, but limited study has found shortened telomere length among those with more perceived stress ⁹³ (Fig. 34-10) and in caregivers, ⁹⁴ and a study suggests an association between pessimism and shorter telomeres as well. ⁹⁵

FIGURE 34-9 Schematic representation of the principal pathways by which chronic stress and affective disorders promote disease. The primary effect of perceived stress on chronic negative emotions is chronic activation of the hypothalamic-pituitary-adrenal (HPA) axis and the sympathetic nervous system (SNS). Widespread peripheral effects derive from this central nervous system activation (see text). In addition, there are direct effects on central nervous system remodeling that may help drive altered behaviors. Another consequence is a heightened physiologic responsivity to acute environmental stimuli. ANS, autonomic nervous system. (From Rozanski A, Blumenthal JA, Davidson KW, et al: The epidemiology, pathophysiology, and management of psychosocial risk factors in cardiac practice: the emerging field of behavioral cardiology. J Am Coll Cardiol 45:637, 2005.)

Finally, extensive research by McEwen and others has established that the brain, itself, is an important end-organ target of chronic stress. Specifically, chronic stress leads to profound primary alterations in brain chemistry and circuitry, including functional remodeling of the amygdala and hippocampus within the brain's limbic system and in the prefrontal cortex. 96 Under conditions of chronic stress, the hippocampus, an essential structure for memory and spatial navigation, manifests functional atrophy of its neural den drites; concomitantly, the amygdala, a region that regulates fear responses and is involved in hormonal, autonomic, and behavioral responses to stress, manifests growth and expansion in its neuronal dendrites (Fig. 34-11). The prefrontal cortex, the master region for executive decision making and regulation of neurohumoral and autonomic responses during stress, also manifests shrinkage of its dendrites during chronic stress. The functional consequences of these central nervous system effects may include impairment in judgment, attention , and memory and an increased vulnerability towards anxiety. Importantly, both the shrinkage in the hippocampus and prefrontal cortex and the hypertrophy of the amygdala are reversible in animal models of stress, and limited human imaging data suggest that this may also be the case in humans. 96

Effects of Depression

Like chronic stress, depression is also associated with widespread systemic effects resulting from activation of the

Prefrontal cortex and hippocampus

FIGURE 34-11 Directional changes in neuronal branching and connectivity under conditions of chronic stress. Chronic stress causes functional atrophy of neuronal dendrites in the medial

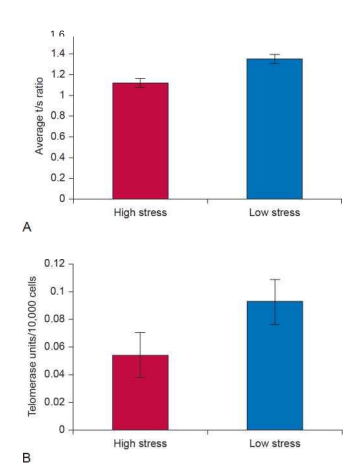
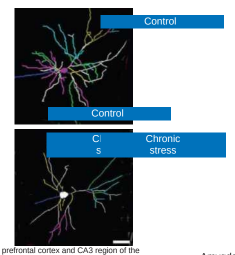


FIGURE 34-10 Comparison of mean telomerase length and telomerase enzyme activity in subjects with high versus low perceived stress. (From Epel ES, Blackburn EH, Lin J, et al: Accelerated telomere shortening in response to life stress. Proc Natl Acad Sci U S A 101:17312, 2004.)



hippocampus and hypertrophy (ie, growth and expansion) in the basolateral amygdala and orbitofrontal cortex. Within animal models, these changes are reversible. (Courtesy of Dr. Bruce McEwen.)

autonomic nervous system and HPA axis. Hypercortisolemia is characteristic in chronic depression, as is heightened stimulation of the sympathetic nervous system, resulting in higher concentrations of circulating plasma norepinephrine and an increase in total body sympathetic activity. As a consequence of the latter, depressed patients commonly manifest autonomic nervous system dysfunction, with decreased heart rate variability, baroreflex dysfunction, increased rest heart rate, and increased QT variability. 97,98 Metabolic abnormalities are also common in depressed patients, including increased insulin resistance, 99 metabolic syndrome, 100 vis ceral fat, 101 and diabetes mellitus, which is increased threefold in depressed patients compared with nondepressed counterparts. 102 The concomitant increase in cortisol and decrease in growth and sex hormone concentrations associated with depression, and possibly induced local inflammatory protein interactions, result in an increased frequency of bone demineralization and osteoporosis. 103 Enhanced platelet activation is also characteristic, apparently resulting from multiple pathophysiological effects of depression. These include elevations in S -thromboglobulin, platelet factor 4, and functional glycoprotein IIb/IIIa receptors; increased responsiveness of platelets to serotonin; and hyperactivity of the 2A receptor 5-hydroxytryptamine transporter transduction system. 97

Substantial data now indicate that depression is also proinflammatory, a result of chronic stimulation of the HPA axis and the sympathetic nervous system, probably acting in concert with various peripheral effects of depression, such as hyperglycemia. Depression is associated with increases in C-reactive protein, fibrinogen, interleukin-6, and tumor necrosis factor and other inflammatory proteins, occurring independently of other CVD risk factors and body mass index. 104 Evidence suggests that the elevation of proinflammatory cytokines contributes to many of the typical somatic symptoms associated with depression, such as fatigue, decreased appetite and weight loss, and sleep and mood disturbances. Another pathophysiological consequence of depression is endothelial dysfunction (Fig. 34-12). 105 This pathophysiological effect may be aided by abnormalities in vascular cell adhesion and proliferation, as manifested by increased levels of intercellular adhesion molecules among depressed subjects. Together, these changes produce a strongly proatherosclerotic environment.

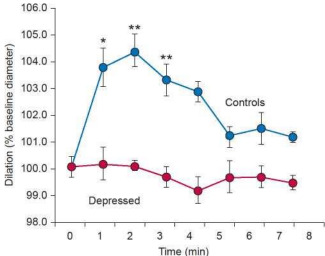
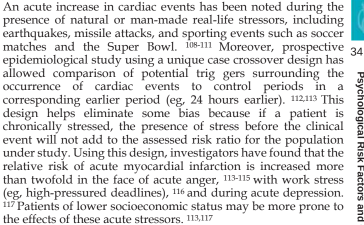


FIGURE 34-12 Comparison of degree of flow-mediated artery dilation after cuff release in 12 depressed patients and 10 healthy controls. Arterial endothelial function is significantly impaired in depressed patients. (From Broadley AJ, Korszun A, Jones CJ, Frenneaux MP: Arterial endothelial function is impaired in treated depression. Heart 88:521, 2002.)

Behavioral Mechanisms

Besides their direct pathophysiological effects, common to all psychosocial factors is strong evidence linking these factors to adverse health behaviors, such as poor diet and overeating, smoking, lack of exercise, poor sleep hygiene, social isolation , and decreased adherence to recommended health habit change. For instance, meta-analyses indicate substantially increased noncompliance with medical treatment regimens among patients with depression ¹⁰⁶ and poor social support. ¹⁰⁷ In the long term, these negative health behaviors both increase the risk for CAD and lead to a vicious circle of a decreased sense of physical and psychological well-being and worsening psychosocial status.

Acute Psychological Stress



The pathophysiological mechanisms that may place cardiac patients at increased risk for myocardial ischemia and cardiac events appear to be multifactorial. Acute psychological stress causes sudden increase in heart rate and often substantial blood pressure elevations, 118 thus increasing myocardial oxygen demand. Simultaneously, among patients with coronary disease, acute psychological stress can cause coronary vasoconstriction, through an endothelium-dependent mechanism, at the sites of coronary stenoses, thus resulting in a concomitant decrease in myocardial oxygen supply. 119 In the laboratory setting, mild psychological stressors can cause transient myocardial perfusion defects 120 and the induction of transient wall motion abnormalities 3; the transient wall motion abnormalities are inducible in approximately half of patients with exerciseinducible myocardial ischemia. Patients who manifest ischemia during mental stress testing are at increased risk of mortality. 121,122 In one study, three different types of mental stressors were compared for their ability to induce regional wall motion abnormalities among CAD patients. Notably, a stressor involving speaking on issues of personal stress was substantially more potent than other mental tasks (the Stroop Word Task and math stress) in inducing myocardial ischemia. When CAD patients are monitored for ischemia with ambulatory electrocardiographic monitoring, ischemia is also often noted during nonexercise stressors, including those of an emotional nature. 123,124 Interestingly , mental stress-induced ischemia is most commonly clinically "silent" (ie, it occurs in the absence of chest pain), and it occurs at a substantially lower double product compared with exercise-induced ischemia. Overall, mental stress-induced ischemia is not common among patients who do not have exercise-induced myocardial ischemia.

Besides these pathophysiological effects, acute mental stress may cause a worsening of endothelial function that may last for hours. ¹²⁵ Pretreatment with metapyrone, a competitive



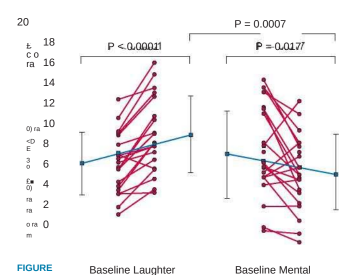
inhibitor of the conversion of 11-deoxycortisol to cortisol, can ablate this effect of acute mental stress. 126 Other work has demonstrated that acute mental stress has the ability to induce platelet activation, 127 to augment indices of arterial wall stiffness, ¹²⁸ and to promote inflammation. ¹²⁹⁻¹³¹ In one unique study that was designed to examine the prospective effects of a real-life stressor on inflammatory and vasoconstrictive mediators, investigators compared the serum from subjects watching World Cup soccer with that from reference groups of patients with acute coronary syndrome and healthy volunteers. 131 Various inflammatory and vasoconstrictor mediators, such as monocyte chemoattractant protein 1 and endothelin 1, were substantially elevated among the subjects who were watching the soccer matches. In addition, an extensive literature has demonstrated the ability of acute mental stress to produce a more proarrhythmic environment and to stimulate arrhythmias, both in the laboratory and in real-life settings. 3,132

Paralleling the general increased interest in positive psychological factors that may buffer against illness, there has also been a recent study in the potential acute beneficial effects of positive moods on physiology. For example, Frederickson and Levenson ¹³³ have demonstrated that whereas negative moods that are induced in the laboratory setting can lengthen the time of hemodynamic recovery from an acute mental stressor, the preinduction of a positive mood can shorten hemodynamic recovery. In other work, films designed to produce laughter and mental stress produced divergent effects on indices of arterial wall stiffness. ¹³⁴ Using a similar design, other investigators have demonstrated divergent effects of viewing laughter versus mental stress-related films on subsequent endothelial function ¹³⁵ (Fig. 34-13). The clinical significance of such findings awaits further study.

Stress Cardiomyopathy

34-13

Acute stress also has the ability to produce an uncommon but increasingly recognized form of acute reversible heart failure, associated with a distinct pattern of acute left ventricular hypocontraction, involving prominent apical akinesis with relative sparing of the base of the left ventricle. ¹³⁶⁻¹³⁸ This



Assessment of brachial artery flow-mediated vasodilation in 20 healthy volunteers at baseline and during viewing of a cinematic film designed to induce laughter (left) and at baseline and during viewing of a film designed to induce mental stress (right). The mean flow-mediated dilation was increased by 22% ± 15% in testing that was performed after inducing laughter and decreased by 35% ± 47% after viewing the mental stress film clips. (From Miller M, Mangano C, Park Y, et al: Impact on cinematic viewing on endothelial function. Heart 92:261, 2006.)

syndrome has been referred to by different names, including takotsubo cardiomyopathy and apical ballooning syndrome. A study has characterized a relatively large group of 130 such patients. ¹³⁹ Predominantly, these patients, as in most series, were women, with stress cardiomyopathy induced by either intensely emotional or physical stress in the majority of these patients. However, some patients had no identifiable trigger, and in a very small number, there was either delayed return of ventricular function or evidence of left ventricular thrombi, raising the issue of screening some patients for anticoagulation.

THE CLINICAL MANAGEMENT OF PSYCHOLOGICAL RISK FACTORS IN CARDIOLOGY PRACTICE

Psychosocial problems such as depression are highly distressing and severely damage quality of life. Moreover, the convincing pathophysiology and epidemiology associated psychosocial risk factors for CAD form a further compelling reason for the development of behavioral interventions to protect against their adverse effects. Because psychosocial problems are commonly concentrated in cardiac patients, screening will provide cardiologists with the opportunity to identify and manage or refer patients with such problems in clinical practice. To date, however, there are no accepted practical guidelines to steer the management of psychosocial problems in clinical practice. Issues to be resolved include a lack of evidence base to suggest optimal therapies; the lack of organized health care systems to aid physicians in such management, particularly as it relates to time constraints that physicians commonly face; and the lack of reimbursement for practicing behavioral interventions. These issues are addressed in this section.

Evidence Base Regarding Behavioral Interventions

To date, there have been only sparse large studies that specifically evaluated the clinical efficacy of stand-alone behavioral interventions in cardiac populations (Table 34-2). The first of these was the Recurrent Coronary Prevention Project study. 140 In this study, 862 individuals were randomized into two groups; 270 received group counseling, and 592 received both group counseling and type A behavioral modification counseling. Another 151 patients served as a non-treated com parison group. During a 4.5-year follow-up, the occurrence of subsequent cardiovascular mortality or nonfatal myocardial infarction was significantly lower in the group that received type A behavioral counseling compared with the other two groups. Subsequently, the Ischemic Heart Disease study demonstrated a reduction in cardiac events among a group of patients who underwent a home-based stress reduction program. 141 Two other large-scale stress management studies were negative, but in neither study did the intervention actually reduce psychological distress. 142,143 In a second analysis of one of these two studies, a significant reduction in cardiac mortality was noted for the subgroup with reduction in distress within the psychological intervention arm of the study. 144

The fifth and most recent large-scale behavioral intervention study was the Enhancing Recovery in Coronary Heart Disease Patients (ENRICHD) trial. ¹⁴⁵ This differed from the prior studies in that it focused on alleviating depression or low perceived social support. This trial randomized 2841 acute post-myocardial infarction patients who had evidence of either depression or low

Investigations	Control Group	Intervention Group	Follow-up	Type of Intervention	Reduced Psychosocial Stress?	Reduced Cardiac Events?
Friedman et al, ∞ 1986	270	592	4.5 years	Type A behavior pattern modification/group counseling	Yes	Yes
Frasure-Smith and Prince,141 1985	229	232	5 years	Home-based nursing intervention	Yes	Yes
Jones and West, 142 1 9 9 6	1155	1159	1 year	Group stress management sessions	No	No
Frasure-Smith et al,143 1997	684	692	1 year	Home-based nursing intervention	No	No*
ENRICHD investigators,145 2 0 03	1243	1238 depression	2.4 years	Cognitive-behavior therapy for	Yes ₄	No*

^{*}Secondary analysis with the intervention groups revealed decreased cardiac events among those patients with reduced stress

either usual medical care or a treatment arm that employs cognitive-behavioral therapy, supplemented by the use of a selective serotonin reuptake inhibitor (SSRI) for severe or unremitting depression. Physicians were allowed to also use SSRIs within the usual care group. A similar frequency of the composite endpoint of all-cause mortality and nonfatal myocardial infarction was observed in both treatment arms after 29 months of follow-up. The results of this trial may have been influenced by a greater than expected reduction in depression and improvement in social support in the usual care group, resulting in only modest difference in psychosocial functioning between both treatment arms by the end of the trial. Also, potential differences may have been obscured by a relatively high use of antidepressants in both treatment arms and a high rate of referral of recruits to myocardial revascularization, occurring in approximately 40% of the patients within 12 weeks of acute myocardial infarction. A later secondary analysis of the ENRICHD trial found that within the intervention arm, those who responded with a reduction in depression had a lower risk for late mortality compared with patients whose depression persisted or worsened. 146

The combined efficacy of psychosocial interventions that involved smaller sample sizes has been assessed in a meta analysis of 36 psychological intervention trials involving 12,851 patients. 147 Overall, these interventions were found to result in only a small reduction in the frequency of nonfatal myocardial reinfarction and no reduction in cardiac mortality or the frequency of subsequent myocardial revascularization procedures. Among the findings emphasized in this meta analysis was the observation that these trials frequently yielded negligible improvement in depression or anxiety levels. In the absence of such an effect, the utility of such studies must be interpreted with caution.

A variety of investigators have attempted to address the paucity of clinical trial evidence on cardiovascular outcomes by examining the effect of psychosocial interventions on intermediary processes that are connected to atherosclerosis or risk for cardiac events. As an example, Blumenthal and colleagues compared the beneficial effects of three management approaches - exercise, stress management, and usual care among three intermediate endpoints in 134 randomized patients (changes in flow-mediated dilation, left ven tricular ejection fraction, and wall motion during radionuclide ventriculography) and in variables reflecting cardiac autonomic control. 148 Both among patients assigned to exercise and stress management, favorable findings in these indices were observed compared with patients assigned to usual medical care. A review of studies that have examined the effects of psychological interventions on such intermediary endpoints is beyond the purview of this chapter, but a practical problem posed by these smaller trials is that they 34 often involve considerably more intense caregiver interactions than is possible in everyday practice.

The end result of the evidence base to date is a lack of certainty as to best practical approaches for management of psychosocial distress in cardiac practice. Nevertheless, in the absence of formal data, there are many insights, practical experiences, and new work in the arena of medical psychology that may guide physicians regarding the management of psychosocial issues in cardiac patients, as discussed here.

Screening for Psychological Distress in Cardiac **Populations**

A recent American Heart Association Science Advisory calls for the routine screening for depression in cardiac practice and suggests that at a minimum, a two-question survey from the Patient Health Questionnaire (PHQ) be applied. 148 The two-item questionnaire includes the following:

Over the past 2 weeks, how often have you been bothered by any of the following problems?

- 1. Little interest or pleasure in doing things.
- 2. Feeling down, depressed, or hopeless.

If the answer to either question is yes, the Advisory suggests that the full nine items of the PHQ be assessed (the PHQ-9; Box 34-2). Patients with high PHQ scores (> 10) should be referred for qualified professional evaluation according to this Advisory.

A number of issues are pertinent relative to this new recommendation. First, to date, there is insufficient study to determine whether the screening for depression, per se, without additional structured follow-up sufficiently impacts on clinical care. Notably, a meta-analysis suggests that the stand-alone use of a depression-screening question has little impact on depression management. 149 In primary care settings, the utility of screening for depression is improved if such screening is tied to some form of collaborative care intervention following screening. 150,151 Second, the American Heart Association Science Advisory did not deal with recommendations for screening for anxiety, which, although substantially comorbid with depression, may occur in its absence and exert a significant effect on morbidity, mortality, health cost, and quality of life. Third, and significantly, despite the widespread emphasis on DSM-IV diagnoses, they largely cover depression, anxiety, and substance abuse disorders; but as reviewed in this chapter, there are a large variety of other psychosocial problems that can exert significant clinical and pathophysiological effects, including chronic stress and social isolation. Whereas there are also short version questionnaires that can be used to screen for these psychosocial factors, the use of questionnaires is probably



Substantial decreases in depression scores were noted in both the control and intervention groups.

30X 34-2 Patient Health Questionnaire-9 (PHQ-9) fo Screening for Depression

Over the past 2 weeks, how often have you been bothered by any of the following problems?

- 1. Little interest or pleasure in doing things.
- 2. Feeling down, depressed, or hopeless.
- 3. Trouble falling asleep, staying asleep, or sleeping too much.
- 4. Feeling tired or having little energy.
- Poor appetite or overeating.
- Feeling bad about yourself, feeling that you are a failure, or feeling that you have let yourself or your family down.
- Trouble concentrating on things such as reading the newspa per or watching television.
- 8. Moving or speaking so slowly that other people could have noticed.
- Thinking that you would be better off dead or that you want to hurt yourself in some way.

Questions are scored: not at all = 0; several days = 1; more than half the days = 2; and nearly every day = 3. Add together the item scores to get a total score for depression severity.

From Lichtman JH, Bigger JT, Blumenthal JA, et al: Depression and coronary heart disease recommendations for screening, referral, and treatment. *Circulation* 118:1768, 2008. Reproduced with permission.



3OX 34-3 Suggested Open-Ended Questions to Screen for Psychosocial Risk Factors

How would you describe your energy level?
How have you been sleeping?
How has your mood been recently?
Do you feel anxious or unduly worried?
Are you under undue pressure at work or at home?
Do you have difficulty unwinding after work or at the end of the day?
Who do you turn to for support?

Modified from Rozanski A, Blumenthal JA, Davidson KW, et al: The epidemiology, pathophysiology, and management of psychosocial risk factors in cardiac practice: the emerging field of behavioral cardiology. *J Am Coll Cardiol* 45:637, 2005.

unnecessary in daily practice. Rather, just as physicians are accustomed to performing a quick oral review of all organ systems in daily practice, a very short series of questions, such as those proposed in Box 34-3, can be used to screen for chronic negative emotions, chronic stress, and somatic complaints that may be stress or emotion related. There is a conception that such investigation is necessarily lengthy and thus impractical, but to the contrary, a practiced clinician can often obtain relevant screening information in a matter of minutes by using simple questions such as those listed in Box 34-3.

Synergy of Managing "Behavioral" and Psychological Risk Factors

There is a common division made between the management of behavioral lifestyle factors associated with CAD, such as smoking, poor diet, and sedentary behavior, and the management of psychological risk factors, but these factors only cluster together. There are reasons that grouping the consideration of these factors is beneficial. Psychological risk factors, such as depression, anxiety, and chronic stress, are

causative factors with respect to poor health behaviors, and they constitute a strong barrier to the successful adherence of recommended health behavior change. 106,107 Equally important, poor health behavior is one of the mechanisms by which psychological risk factors cause or worsen CAD, and favorable behavioral change is one means of ameliorating some aspects of psychological distress.

As an example, exercise represents a behavioral intervention that can be used to ameliorate psychological distress, as shown by cross-sectional and randomized controlled trials demonstrating lower depression among those who exercise. ^{152,153} In other work, Blumenthal and coworkers randomly assigned 156 depressed subjects to exercise, antidepressant medication, or both. ¹⁵⁴ Exercise was just as effective as ser traline hydrochloride in reducing depressive symptoms in this cohort, but the study did not contain a control group.

A subsequent randomized controlled trial compared four therapies: supervised group exercise, home-based exercise, antidepressant medication, and placebo groups. ¹⁵⁵ Comparable reductions in depression were again observed with exercise and antidepressant medication, with a lesser but beneficial effect associated with placebo medication. Further, examination of the effects of exercise within the ENRICHD study found that it was associated with both a significant reduction in depression and a 30% reduction in mortality. ¹⁵⁶ A cardiac rehabilitation program is a common clinical setting in which psychosocial interventions have been combined with exercise training and standard behavioral risk factor modification.

In a recent meta-analysis of 23 randomized controlled trials that involved cardiac rehabilitation, Linden and colleagues ¹⁵⁷ observed a reduction in all-cause mortality at 2-year follow-up for those studies incorporating psychosocial interventions compared with those that did not (odds ratio of 0.72; 95% CI, 0.56-0.94). Notably, paralleling results seen among studies involving stand-alone psychosocial interventions, reduction in mortality was significant only among patients in whom there was successful reduction of psychological distress within this meta-analysis.

New Potential Paradigms for Psychosocial Interventions

Potential advances in the psychosocial management of cardiac patients can be inhibited when strict boundaries are maintained between what cardiologists and behavioral specialists manage. New insights into the determinants and means of increasing human motivation, the bidirectional relationship between thoughts and emotions, and the potential application of positive psychology practices represent new opportunities to develop more optimal and potentially efficient interventions in cardiac practice.

An example of how cardiologists might think about stress, moods, emotions, and health behaviors in an integrative way stems from work conducted by Thayer and colleagues. ¹⁵⁸ In a series of experimental subjects, they assessed the relationship between four parameters: feelings of tension, energy levels, moods, and behaviors. The paradigm that evolved from their work revolves around four energy-tension states, as illustrated in Figure 34-14. These investigators noted that when subjects reported low tension, whether in association with high energy (ie, "calm energy") or low energy (ie, "calm tiredness"), corresponding mood states were generally positive. By contrast, when subjects reported high tension, moods varied according to subjects' corresponding energy levels; "tense tiredness" was associated with a high frequency of negative moods, but "tense energy" was not.

These findings offer insights into a potential management paradigm, involving four categories of intervention, for the amelioration of psychosocial risk factors in cardiac practice.



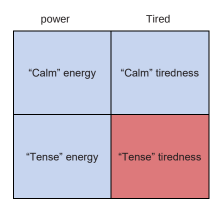


FIGURE 34-14 Thayer's proposed model concerning characterization of subjects' momentary states according to energy and tension levels. The majority of negative moods in this experimental work occurred in subjects when they reported both high tension and low energy (ie, "tense tiredness"). (From Thayer RE, Newman R, McClain TM: Self-regulation of mood: strategies for changing a bad mood, raising energy and reducing tension. J Pers Soc Psychol 67:910, 1994.)

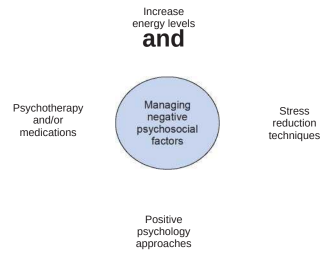


FIGURE 34-15 Potential methods for management of negative psychosocial interventions in clinical practice (see text).

Whereas these categories are listed as separate approaches in Figure 34-15, in fact there is considerable overlap and potential synergy among these approaches. One approach is to focus subjects on managing their energy. This can be beneficial because when patients are under prolonged stress, there is often a tendency to become more self-centered and to feel greater urgency to manage stress, even though at such times patients may not be at their best for problem solving. 68 The work of Thayer implies that when patients lift their energy, their mood may also lift, thus providing them with greater self-efficacy for problem solving afterwards. Patients may benefit from advice on how to increase their energy effectively . A common quick fix for patients when they are feeling tense-tired is the ingestion of unhealthy energy-dense foods. 158,159 Rather, physicians can suggest such energy boosting activities as short bursts of exercise activity, switching from draining work to inspiring work, taking naps, or listening to inspiring music.

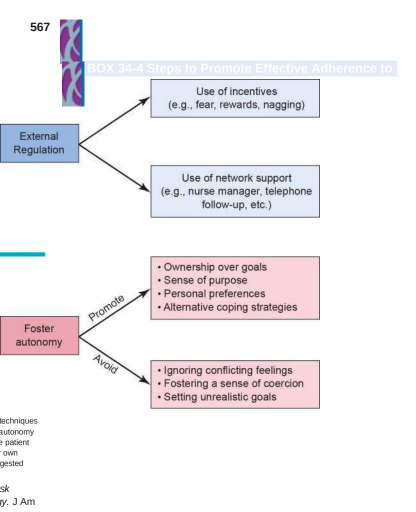
Alternatively, methods that are designed to physically and emotionally unwind or reduce stress may be useful, including such practices as simple breathing exercises and the application of mindful meditation. Practices such as yoga and Tai Chi represent approaches that can both increase energy and reduce stress when they are employed over time. There is increasing evidence that such techniques have beneficial physiological effects. ¹⁶⁰⁻¹⁶²

When patients are tense-tired or moody, there is a tendency to become more cognitively inflexible, impatient, and less creative. ⁶⁸ A behavioral approach that teaches patients to take their negative thoughts, moods, and problems less seriously or to turn away from these thoughts to positive pursuits that generate positive emotions is becoming commonplace among practitioners of positive psychology and might be helpful to cardiac patients as well. As an example of this approach, a couple under marital stress may choose to focus on their specific complaints under standard marital therapy. In this alternative approach, a positive psychology practitioner may have the couple focus and dwell on the positive aspects of their relationship during counseling sessions, with the intention of raising emotional well-being, with enhanced problem solving evolving as a secondary outcome of that effort. To date, there is a dearth of studies regarding the potential effects of positive psychological interventions on clinical outcomes, but the beneficial effects of positive interventions on some mediators of physiology and central nervous system function have been noteworthy. 163,164

When stress, anxiety, or mood disorders become more severe, referral of patients to trained mental health professionals for formal psychotherapy should be considered. Some sources of stress and the health behaviors they induce, such as overeating and sedentary behavior, poor social support, loneliness, overwork, and life imbalance (sometimes leading to unduly severe voluntary sleep restriction), may be ame nable to physician advice or referral to community or Internet resources. Notably, many of these psychosocial problems do not necessarily fall within the traditional division of what physicians versus mental health care specialists manage.

The Use of Psychopharmacological Medications

When depression or anxiety is sufficiently severe, patients are candidates for the use of pharmacological agents or some form of professional psychotherapeutic counseling, such as cognitivebehavioral therapy or problem-solving therapy. Cardiologists do not commonly manage depression, but they may either refer patients directly to psychiatrists for psychiatric evaluation or work with general practitioners who are accustomed to treating depression or collaborating with psychiatrists for mental health care. In either case, it behooves cardiologists to be well acquainted with the medical side effects psychopharmacological agents and potential drug interactions. In recent years, SSRIs have become the preferred medication for the treatment of depression in cardiac patients, and they are also used as a first-line medication for the treatment of significant anxiety. SSRI medication (sertraline) was found to have a very good safety profile for use in post-myocardial infarction patients in the SADHART study, 165 but recent studies have suggested that the use of SSRIs is not necessarily benign. A 5.9-year followup of 136,293 women from the Women's Health Initiative found that SSRIs were not associated with an increased risk of cardiac events, but there was an increased risk for all-cause mortality and hemorrhagic and fatal stroke. 166 However, the increase in absolute event rate with SSRI use during this follow-up was still low. In another large follow-up of 63,469 women from the Women's Health Study, Whang and associates 167 found that major depression and antidepressant use were associated with an increased cardiac event rate, and antidepressant use alone was associated with an increased event rate for sudden cardiac arrest. It is not clear whether this effect merely signaled the presence of greater underlying



Behavioral Suggestions

- Use clear and effective communication and make recommendations as specific and simple as possible.
- Schedule follow-up visits to check adherence, especially during the early practice phase, as opposed to the later, more ingrained habit phase.
- Provide a motivating rationale for the patient's treatment regimen, with consideration of explanations that benefit the patient's level of "health literacy."
- Follow key oral suggestions with written ones to reinforce the cardiologist's message.
- Begin with "micro" goals for patients who are resistant to behavior change or who have fewer available personal resources.
- 6. Help patients establish realistic goals and expectations.
- Involve patients in tailoring behavioral suggestions that reflect their autonomous desires and tendencies rather than just dictating change.
- 8. Provide positive feedback.
- 9. Openly and candidly explore potential patient barriers to adherence (such as lack of personal motivation, time, family support, facilities, or knowledge; fears; job, home, or other pressures; and cultural issues) and assist patients with problem solving and developing strategies (eg, self monitoring approaches, written agreements, and relapse prevention) at the time of recommendations.
- 10. Make use of "implementation intentions."
- 11. Help patients to become cognizant of environmental stimuli that may inadvertently prime negative health behaviors unconsciously.
- Refer patients with poor structural or functional social support to programs or activities that will enhance adherence by providing social support.

depression among those taking antidepressant medication or represented an important side effect of SSRI use. Such recent observations point out the need for further prospective study regarding the use of SSRIs in cardiac patients, both to study their potential benefits in promoting survival, which to date has been poorly studied, and to better assess potential medical side effects of SSRI use in cardiac patients. Still, these medications enjoy a substantially improved patient profile compared with various prior generations of antidepressant agents, such as tricyclic depressants.

The Psychological Management of Patient Adherence

Behavioral risk factors for CAD are well established, including sedentary behavior, smoking, overeating, and poor nutritional diets. Depression, anxiety, poor social support, and other psychosocial risk factors offer a powerful barrier to patients' attempts to modify these adverse health behaviors as well as to medication adherence. Research indicates that understanding of psychological principles can increase patients' internal motivation and help them plan and maintain adherence to health behaviors. For instance, patients become more motivated and they relapse less often when they formulate health goals that are more consonant with their own autonomously endorsed values or patterns of behavior, rather than just being told by their physicians what to do 97 (Fig. 34-16). Increasingly, it is also recognized that patients are unconsciously or "mindlessly" cued into behav iors by their external environment. 168,169 Along these lines, physicians can advise patients about environmental factors that induce "mindless eating" 170 or promote stress. The

Modified from Rozanski A, Blumenthal JA, Davidson KW, et al: The epidemiology, pathophysiology, and management of psychosocial risk factors in cardiac practice: the emerging field of behavioral cardiology. *J Am Coll Cardiol* 45:637, 2005.

tendency to behave unconsciously can also be positively manipulated to promote healthy behaviors through the formulation of "implementation intentions," as developed by Gollwitzer. ¹⁷¹ Implementation intentions involve patients' declaration that when it is X, I will do Y. For instance, "when I finish work at 6 PM at night, I will drive to the gym before coming home." Although this formulation seems simplistic, meta-analysis indicates that it can be effective in fostering behavioral change. ¹⁷² Approaches to augment patients' adherence to recommended health behaviors are listed in Box 34-4.

Potential Practice Model for Psychosocial Interventions in Cardiac Practice

The management of both behavioral and psychological problems varies widely in complexity and in terms of time and resource demands. A potential stepped care model for integration of psychosocial interventions into medical care is shown in Figure 34-17. Patients with milder forms of psycho social distress and with a tendency to be highly adherent to medical suggestions can be readily managed within routine physician practice. ¹⁷³ Physicians can enhance their management of patients by being aware of principles designed to promote the adherence of patients to suggested behavioral change. ⁹⁷ Whereas physician time is limited, physicians can complement their management of patients by suggesting to them community-based exercise and social programs and

STEPPED CARE BEHAVIORAL MANAGEMENT FOR CLINICAL PRACTICE

Refer to behavior specialists

Increasing psychosocial or adherence risk

Physician management

FIGURE 34-17 A potential model for stepped psychosocial interventions in clinical cardiac practice. Routine counseling and psychosocial interventions may be addressed at the physician level. Somewhat increased difficulty in adherence to medical regimens or psychosocial stress may benefit from adjunctive care, monitoring, or feedback from office staff (see text). Patients with significant psychosocial distress (eg, depression) or poor adherence difficulty may benefit from additional referral to behavioral health specialists or programs. 3

Internet-based programs that can support dietary and other behavioral changes. The range of such programs varies widely within local communities. For patients with somewhat increased psychosocial distress or with greater difficulty in adhering to behavioral change, added instruction, support, monitoring, or feedback might be accomplished by use of ancillary office staff.

Added support is often quite helpful in the early stages of requested behavioral changes because unlike habits, which are automatic and effortless, new behavioral practices are intentional and effortful. Patients who are poorly adherent or resistant to recommended behavioral change or manifest a high level of psychological distress can be referred to appropriate health care professionals, according to the nature of the patient's problems. Prospective work is needed to test the potential of this stepped care model.

CONCLUSION

Epidemiological studies have been consistent in demonstrating a strong relationship between various psychosocial risk factors and heart disease. These factors include chronic negative emotions, such as depression and various forms of anxiety; negative cognitive patterns, such as pessimism; chronic stress; and the lack of certain basic psychological needs, such as poor social support. In general, a strong dose response association exists among many psychosocial risk factors and CAD outcomes. Importantly, new evidence indicates that positive psychological factors and life experiences can benefit physiology and improve longevity. Great strides have been made into understanding of the pathophysiology linking psychosocial risk factors to CAD, involving chronic activation of the autonomic nervous system and the hypothalamic-pituitary axis, leading to remodeling of central nervous system structures, physiological hyperreactivity, apparent accelerated aging, and negative metabolic, coagula tion and immune effects that serve to promote accelerated atherosclerosis and increase the risk for cardiac events. Results of psychosocial intervention trials have resulted in mixed data, but new paradigms and developments within medical psychology offer new potential means to improve the efficacy of psychosocial interventions in cardiac patients.

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Office staff

assistance

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CHAPTER 35

The Role of Treatment Adherence in Cardiac Risk Factor Modification

Thomas M. Maddox and P. Michael Ho

KEY POINTS

- Treatment adherence is essential for the optimal control of cardiac risk factors.
- Poor adherence, with subsequent adverse cardiac outcomes, is evident in hypertension, hyperlipidemia, diabetes, smoking cessation, and weight loss.
- Elements associated with poor adherence include social and economic factors, health care system and team-related factors, condition-related factors, therapy-related factors, and patient-related factors.
- Medication simplification and team-based approaches to identification and treatment of cardiac risk factors have shown promise in improving treatment adherence.
- Future approaches to improvement of treatment adherence may include moving to a home-based model of care, activating patients to participate in their care, and employing information technology to identify nonadherence and to initiate interventions.

Drugs don't work in patients who don't take them.

C. Everett Koop

For patients with coronary artery disease (CAD) or at risk for CAD, optimal control of cardiac risk factors, such as hypertension, hyperlipidemia, and smoking, is one of the most effective methods to reduce subsequent cardiac morbidity and mortality. For this control to be achieved, a variety of therapeutic options, both pharmacological and nonpharmacological, are available to patients and their health care providers. However, none of these therapeutic options can provide optimal cardiac risk factor control without diligent adherence to their use. Indeed, treatment adherence serves as the "final common pathway" for risk factor treatment and has been characterized as the "key mediator between medical practice and patient outcomes." 1 Despite this essential and somewhat obvious role in risk factor optimization, treatment adherence has been a relatively neglected area of research because of difficulties in both characterizing the factors that drive adherence and designing effective interventions for improvement. Thus, a critical step in optimizing cardiac risk factors in patients with CAD is understanding and improving treatment adherence.

Adherence to treatments of cardiac risk factors is difficult for a variety of reasons. Many risk factors, such as hypertension and hyperlipidemia, are chronic, life-long conditions that require daily administration of treatments, including both pharmacological agents and lifestyle choices. Furthermore, these conditions are largely asymptomatic. Thus, an important motivator for treatment – relief from bothersome symptoms – is largely absent in the management of cardiac risk factors. In fact, the treatments themselves may cause symptoms via side effects, thus adding another disincentive for adherence among affected patients. Finally, adherence is also influenced by a variety of patient, provider, and health care system characteristics.

This chapter reviews the current definitions of treatment adherence, methods of its measurement, and current gaps in the treatment of cardiac risk factors. Although both lifestyle and pharmacological treatments are important for optimal risk factor control, the bulk of the research has focused on adherence patterns to pharmacological therapies and thus comprises the majority of the review. Barriers to treatment adherence, including those at the patient, provider, and health care system levels, are reviewed. Interventions to improve adherence are also reviewed for both their relative effectiveness and potential dissemination. Finally, future directions for adherence research and interventions are presented.

ADHERENCE DEFINED

Treatment adherence is defined as the extent to which patients take treatments (medications or lifestyle modifications) as prescribed by their health care providers. ² A variety of terms have been used to describe treatment adherence in the past, such as compliance, concordance, and fidelity, but these terms have been largely abandoned because of the pejorative nature underlying their use. 3,4 Adherence is usually measured on a continuous scale as a percentage calculated by the number of treatments actually taken relative to the number of treatments prescribed. 5 For example, a patient who is prescribed 30 pills of an anti hypertensive medication and actually takes only 20 pills is 67% adherent to the prescribed regimen. The adherence scale can range from 0% to > 100% (because some patients can take more than their prescribed treatments). 5 Truly perfect adherence, with fidelity to precise timing of each dose of treatment, is rare, occurring in only one sixth of patients in one survey. 6,7 Furthermore, many of the treatments for cardiac risk factors (eg, statin therapy for hyper lipidemia) may not require such precision for optimal control to be achieved. Accordingly, most studies on adherence have used a dichotomous cutoff of > 80% to define good versus poor treatment adherence. ⁵

A related but separate concept to treatment adherence is treatment persistence.

Persistence is the duration of time that a patient remains on delays in medication administration, rather than outright treatment from the initiation to discontinuation of a prescribed discontinuation of treatments. Urquhart and colleagues char due to side effects or provider-initiated discontinuation.

variety of ways. In general, measurement can be direct or indirect, behavior, taking almost all of their doses at their prescribed with tradeoffs made between precision and ease of measurement frequency. One sixth took all of the prescribed doses but had some for each type of method. Direct methods of adherence timing irregularity. One sixth had an occasional missed dose, along measurement include directly observed therapy and measurement with some timing irregularity. One sixth of patients took "drug of the blood concentration of a medication or its metabolite. Both of holidays," or several consecutive missed doses, three or four times these techniques, although accurate, require substantial time and a year. One sixth took drug holidays monthly or more often, and expense and are thus rarely employed. Indirect methods of the final group of patients took few or no doses, although they adherence measurement include patient self-report of treatment would often report good medication adherence. 6,7,13 Consistent adherence, pill counting, assessment of medication refill rates, with this desire by patients to exhibit good adherence behavior, assessment of a clinical response to a prescribed therapy, and Feinstein and colleagues 14 demonstrated that many patients 571 medication electronic monitoring systems (MEMS) that track the improve adherence in the 5 days preceding a visit to the physician, frequency and timing of opening of the pill containers. In general, a phenomenon dubbed "white coat adherence" by the investigators. self-report and pill counts are not considered reliable methods of assessment, in part because of the social desirability bias that occurs health outcomes and increased health care costs. A variety of when patients exaggerate their behavior in an effort to please their studies have demonstrated that patients with poor adherence have health care providers. This bias also affects the ascertainment of higher all-cause mortality and cardiovascular event rates compared patients' adherence to prescribed lifestyle modifications. On the to those with good adherence. 15-18 In addition, two studies have other hand, medication refill rate assessment is a reliable method of demonstrated that among medication-related hospitalizations in adherence assessment, especially in closed health care systems in the United States, between 39% and 69% are due to nonadherence. which patients receive their medications from a single pharmacy 19,20 Specific studies examining patients with chronic medical source. MEMS assessments are also considered reliable, although conditions such as hypertension and hyperlipidemia have found their use is expensive and logistically complex. Finally, use of a associations between nonadherence and worse treatment outcomes combination of adherence measurement methods can also be a , higher hospitalization rates, and increased health care costs. 21,22 valuable and reliable way to determine adherence patterns. ²

PATTERNS OF **NONADHERENCE** AND **ASSOCIATED OUTCOMES**

Research into adherence patterns for cardiac risk factor treatments has uncovered significant gaps. As previously noted, adherence is generally worse among those patients who have chronic asymptomatic conditions, such as hypertension and hyperlipidemia, than among those with acute symptomatic NONADHERENCE IN HYPERTENSION^{33 34} illnesses. Even among those patients participating in clinical trials, which are optimized for medication provision and follow-up and generally have high adherence rates, adherence rates for chronic conditions average only 43% to 78%. 8-10 In addition, several studies have documented that both adherence to and persistence with chronic treatments drop precipitously after the first 6 months of therapy. 9,11,12

Most deviations from adherent behavior are omissions of or

therapy. 3 Unlike adherence, persistence patterns not only can acterized specific patterns of medication-taking behavior, using describe patient adherence actions but also can identify other MEMS, among patients initiated on chronic preventive therapies. reasons for therapy discontinuation, such as intolerance of therapy They found that patients exhibited six types of medication-taking behavior, with roughly even allocation of patients to each group. Treatment adherence and persistence can be measured in a One sixth of patients exhibited nearly perfect medication-taking

> Predictably, these gaps in treatment adherence lead to worse McCombs and colleagues demonstrated that patients with hypertension who interrupted or terminated their blood pressure treatment accumulated an additional \$873 in health care costs (primarily hospitalization related) during the following year. ²³ This finding was supported by an analysis of the UK-based MediPlus data base that demonstrated an association between the discontinuation of antihypertensive treatments and increased hospital and general physician costs. 24

Hypertension is a main risk factor for the development of both CAD and cerebrovascular disease and a major contributor to worldwide cardiovascular mortality. ²⁵ Uncontrolled hypertension among CAD patients is associated with recurrent cardiovascular events, including death, myocardial infarction, and stroke. ²⁶ Fortunately, a wide variety of treatments exist to achieve blood pressure control and to minimize the risk of these adverse events. Currently, more than

34 classes of antihypertension medications are available, and achieving a sustained decrease in blood pressure of 12 mm Hg with one or more of these treatments can prevent one death for every 11 patients treated. ²⁷

Despite this availability of treatment and evidence of its benefit, blood pressure control for a large number of hypertensive patients remains suboptimal, and nonadherence to treatment is a major factor. Several studies have demonstrated that less than 50% of CAD patients with hypertension have their blood pressure at recommended levels. ^{28,29} Nonadherence is a major contributor to this lack of control and, in one study, appeared to contribute to uncontrolled blood pressure more than the lack of adequate prescription of hypertension treatments. 30,31 Furthermore, patients with greater than 80% adherence rates demonstrated improved blood pressure control relative to those patients with less than 50% adherence rates. Treatment adherence to hypertension medications is especially problematic given the asymptomatic nature of the condition, the common occurrence of side effects associated with hypertension treatments, and the daily commitment required for proper medication adherence. ² These factors all lead to difficulty with adherence. Haynes and colleagues 32 illustrated these difficulties by finding that more than 50% of newly diagnosed hypertensive patients with poor blood pressure control had problems with treatment adherence.

Nonadherence to hypertension treatments with its accompanying uncontrolled blood pressure leads to adverse outcomes. Maronde and colleagues 33 found that Medicare patients who were rehospitalized had significantly higher rates of antihypertension medication nonadherence than a comparable group of nonhospitalized Medicare patients. Similarly, Psaty and colleagues 34 found that patients who were recently nonadherent to beta blocker therapy for their

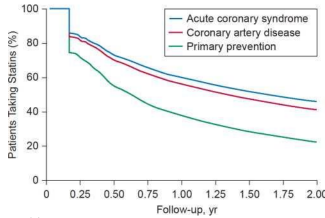
hypertension demonstrated a 4.5-fold increased risk for coronary heart disease complications.

NONADHERENCE IN HYPERLIPIDEMIA

Among CAD patients, lipid-lowering therapies, especially statin medications, can significantly decrease the risk of recurrent cardiac events and mortality. For example, the Heart Protection Study evaluated the effects of simvastatin among CAD, diabetic, or treated hypertensive patients and found an 18% reduction in cardiac-related deaths and a 24% reduction in major cardiovascular events. 35 Similarly, the PROVE-IT TIMI 22 trial compared differing intensities of statin therapies (pravastatin versus atorvastatin) among acute coronary syndrome patients. After 2 years, the atorvastatin patients had lower low-density lipoprotein levels and a 16% reduction in the primary endpoint (death from any cause, No. at Risk myocardial infarction, unstable angina requiring rehospitalization, revascularization, or stroke). 36 Meta-analyses of statin therapy echo these benefits. One involving approximately Primary prevention 25,000 subjects from 34 trials found that 4 years of cholesterol- FIGURE 35-1 Survival curves for lowering therapy would prevent one death, one coronary heart curves are based on a Cox proportional hazards model adjusted for covariates. The median followdisease death, and one cardiovascular death for every 110, 96, and 117 patients treated, respectively. ³⁷ In addition, a survival benefit primary prevention, and 303 days overall. (From Jackevicius CA, Mamdani M, Tu JV: was seen in studies in which more than 50% of patients were Adherence with statin therapy in elderly patients with and without acute myocardial infarction survivors, had a total cholesterol reduction coronary syndromes. JAMA 288:462, 2002.) of more than 10%, and were treated for at least 4 years. In addition, statins produced a greater reduction in the odds of death than other lipid-lowering therapies.

Despite these clear benefits of statin use, nonadherence is a significant problem. Benner and colleagues 38 found that more than half of patients started on statin medications will discontinue the medication within 6 months. Jackevicius and colleagues 35 noted a similar gap in a study of statin adherence patterns among elderly patients. They found that the 2-year rate of statin adherence among this cohort was less than 50%. Furthermore, they demonstrated differential rates of adherence based on the presence of known CAD (Fig. 35-1).

> SURVIVAL CURVES FOR ADHERENCE WITH STATINS IN 3 COHORTS



requiring Acute coronary syndrome 22379 16312 12901 10662 Coronary artery disease 36106 25416 19558 15823 13094 85020 47685 33564 26401 21602

adherence with statins in three cohorts. All up was 494 days for acute coronary syndrome, 430 days for coronary artery disease, 235 days for

Diabetes is another major cardiac risk factor among CAD patients. Diabetic patients are at higher risk for developing CAD and dying from it. 42 In addition, concurrent cardiac risk factors (eg, hypertension, smoking) are more prevalent among diabetic patients and appear to exert worse effects in the setting of poor glycemic control. Diabetes is a complex disorder that requires constant attention to diet, exercise, glucose monitoring, and medication to achieve good glycemic control. 3 Glycemic control in diabetes plays an important role in reducing microvascular disease, such as nephropathy, retinopathy, and neuropathy. 43-45 In addition, a 17-year study of type 1 diabetics indicated that tight glycemic control reduces the risk of cardiac events by 42%. 46 In contrast, tight glycemic control among type 2 diabetics has not demonstrated conclusive benefits for the reduction of cardiac events and may be associated with harm in some cases. 47 Despite this uncertainty, multifactorial cardiac risk factor control, including glycemic control, is an important element of effective prevention of cardiac events. For example, the Steno-1 trial demonstrated the benefit in achieving multiple risk factor control among diabetics. 48 Patients were randomized to usual care or an intensive treatment regimen that included both behavioral and pharmacological treatments targeting hypertension , hyperglycemia, dyslipidemia, microalbuminuria, and secondary prevention of CAD events with aspirin. After 7.8 years of follow-up, patients receiving the intensive treatment experienced a 53% reduction in cardiovascular disease. Moreover, these benefits were sustained, with the patients receiving intensive treatment experiencing a 57% reduction in cardiovascular deaths during the following 4 years after the cessation of the trial.

As with the other cardiac risk factors requiring prolonged chronic treatments, treatment adherence is a substantial problem among diabetics. Among patients receiving oral hypoglycemic therapies, adherence rates range widely from 36% to 93%. 3 Boccuzzi and colleagues 49 profiled adherence rates among patients taking oral antidiabetic medications and found that 12 months after initiation of these therapies, adherence to metformin was 60%; to sulfonylureas, 56%; to repaglinide, 48%; and to a -glucosidase, 31%. Adherence

Although patients who had experienced acute coronary syndrome had higher 2-year adherence rates than those with chronic CAD or receiving statins for primary prevention (40% for acute coronary syndrome, 36% for chronic CAD, and 25% for primary prevention), all patient groups demonstrated significant adherence gaps.

Nonadherence to statins is significantly associated with adverse outcomes. Wei and colleagues 39 found that post myocardial infarction patients who had at least 80% adherence rates to statin therapy experienced relative risk reductions of 81% for recurrent myocardial infarction and 53% for all-cause mortality, but those patients who had adherence rates < 80% had no significant risk reduction in either recurrent cardiac events or mortality. The Lescol Intervention Prevention Study supported these findings by demonstrating that post-myocardial infarction patients who discontinued their statin medications experienced a twofold increase in major adverse cardiovascular events. 40 These benefits of adherence appear to be medication specific, as demonstrated in an elegant study by Rasmussen and colleagues. 41 They found that adherence to statins exhibited a "dose response" relationship with mortality, with higher adherence corresponding to decreased mortality. A similar relationship between calcium channel blockers and mortality among the same cohort was not seen, suggesting that the adherence mortality relationship was statin specific.

NONADHERENCE IN DIABETES

demonstrated that only 63% of insulin doses were taken as replacement therapies, the highest adherence rates were with the prescribed. 3

Nonadherence to diabetic treatments is further complicated by the common occurrence of other cardiac risk factors and comorbidities, many of which require multiple additional medications. Piette and colleagues 50 demonstrated that 50% of US diabetic patients were receiving at least seven medications, including at least two glucose-lowering medications . Adherence rates are adversely affected by this increased number of medications. Rubin ⁵¹ found that rates of adherence to polytherapy versus monotherapy are 10% to 20% lower among diabetic patients. In addition, treatments that require multiple administrations during the day also adversely affect adherence. Paes and colleagues ⁵² found that the adherence rates were 79% for once-daily dosing of

medications, 66% for twice-daily dosing, and 38% for three-timesdaily dosing.

Predictably, nonadherence to antidiabetic medications is linked to adverse outcomes. Pladevall and colleagues 53 illustrated that 10% increases in nonadherence to antidiabetic treatments were associated with 0.14% increases in hemoglobin A1c. Lau and Nau 54 demonstrated that patients who were nonadherent to their risk due to obesity is confounded by the common presence of diabetic regimen were more likely to be hospitalized. In contrast, concurrent cardiac risk factors, obesity appears to be an increased medication adherence to antidiabetic medications (at a independent risk factor for CAD, probably by contributing to threshold rate of 60% +) resulted in decreased medical care costs, insulin resistance, hyper tension, lipid abnormalities, left although overall costs were not reduced because medication costs ventricular hypertrophy, endothelial dysfunction, and obstructive offset these savings. 55 Similarly, Balkrishnan and colleagues 56 demonstrated that 10% increases in medication possession ratios controlled or even eliminated with weight loss. Although no were associated with 8.6% to 28.9% decreases in annual health care randomized controlled clinical trials have linked weight loss to costs.

NON-ADHERENCE IN SMOKING

Smoking is a major cardiac risk factor and remains a significant public health issue. In 2007, 19.8% of US adults were active smokers. 57 Smoking increases all-cause and cardiovascular mortality, and among patients with CAD, it increases the risk of reinfarction and mortality. ^{58,59} Quitting smoking, in turn, reduces all-cause mortality by 36% in CAD patients, even in long-term smokers. 60 In addition, there is some evidence that public health policies such as smoking bans can have salutary effects. A small study in Colorado demonstrated a 27% reduction in myocardial infarction hospitalizations after the institution of a smoking ban. 61 A similar study in Scotland demonstrated a 14% to 21% reduction in acute coronary syndrome admissions during the 10 months after a smoking ban. 62

Quitting smoking is a difficult process, with high relapse rates. For example, in a study of long-term abstinence rates among smokers who tried to quit without any behavioral or pharmacological assistance, only 5% to 7% were abstinent at 1 year. 63 Accordingly, a variety of both behavioral and pharmacological interventions have been developed to assist with smoking cessation. Behavioral interventions include in-person counseling, remote (phone, Web) counseling, group counseling, and financial (eg, patch, gum, lozenge, inhaler, nasal spray), bupropion, and varenicline.

Despite the wide variety of treatments, the rates of adherence to these programs, measured as abstinence from smoking after completion of the treatment, remain remarkably low. For example, quit rates for in-person clinical counseling are 12.3% at 3 months and 6.5% at 12 months 64; for group programs, 20% at 1 year 65; differing and for remote counselling, 10% at 1 year. 66 Financial incentive programs resulted in quit rates of 14.7% at 9 to 12 months and 9.4% at 15 to 18 months. 67 Quit rates with pharmacological therapies are better than with behavioral therapies, but the absolute rates of

rates with insulin therapy were not much better; one study adherence and smoking abstinence remain low. Among nicotine patch, but 12-week abstinence rates were similar for all nicotine replacement modalities (gum, 20%; patch, 21%; spray, 24%; inhaler, 24%). 68 Bupropion resulted in quit rates of 44% (compared with 19% with placebo) at 7 weeks and 23% (compared with 12%) at 1 year. 69 Longer term bupropion (administered for 52 weeks versus 7 weeks) had better short term abstinence rates (47% versus 37%) at 16 weeks after cessation of therapy but similar rates of abstinence to short term bupropion at 2 years (41% versus 40%). Varenicline demonstrated abstinence rates of 44% 4 weeks after medication cessation, compared with 30% with bupropion and 18% with placebo. At 1 year, abstinence rates were 23% with varenicline, 16% with bupropion, and 9% with placebo. 70,71

NONADHERENCE IN OBESITY

Obesity is a growing problem in the United States and worldwide and serves as a risk factor for cardiac disease. The _ | NHANES survey from 1988 to 1991 demonstrated that 36% of surveyed patients were obese. 72 Obesity is associated with coronary disease; Bogers and colleagues 73 showed a 29% increase in CAD with each 5-unit increase in body mass o, index. Although the direct cardiac sleep apnea. 74 Most if not all of these risk factors can be better decreased mortality or CAD, a large observational study found that overweight women who lost more than 20 pounds had a 25% decrease in mortality, CAD, and cancer mortality. Among the subset of women with CAD or heart failure, any weight loss was associated with a 10% reduction in CAD and a 20% reduction in allcause mortality. 75

Similar to smoking cessation, weight loss is difficult to achieve. A variety of weight loss strategies exist and include behavioral modification, dietary therapy, exercise, drug therapy, liposuction, and bariatric surgery. Despite these options, though, adherence to weight loss programs, measured by achievement and maintenance of weight loss, is low.

Behavioral modification strategies include self-monitoring, control of the stimuli that activate eating, slowing down eating, goal setting, behavioral contracting and reinforcement, nutrition education, modification of physical activity, social support, and cognitive restructuring. These techniques are usually paired with dietary weight loss programs, and the combination has been demonstrated to increase weight loss by 7.7 kg at 12 months. 76 In addition, ongoing behavioral modification strategies may be useful in maintaining weight loss. Wing and colleagues 77 showed that those patients who achieved a mean weight loss of 19 kg in the previous year had a lower amount of weight re-gain with in-person support (face-to-face support resulted in weight gain of 2.9 kg; Internet based support, 4.7 kg; and newsletter only, 4.9 kg).

Commercial diet programs, which rely primarily on calorie incentives. Pharmacological treatments for quitting smoking restriction, have demonstrated efficacy in weight loss. Both Weight include nicotine replacement therapies in a variety of formulations Watchers and the Jenny Craig programs have been studied. Weight Watchers resulted in the loss of 5.3% of baseline weight compared with 1.5% in the placebo group at 1 year. ⁷⁸ At 2 years, both groups had regained weight, but the Weight Watchers participants gained back less. Similarly, Jenny Craig participants lost 7.1% of their baseline weight versus 0.7% in the placebo group at 1 year. ⁷⁹ In addition to overall calorie restriction, a variety of diets with



macronutrient composition have been promoted, but data have been conflicting about the superiority of one over the other. For example, a comparison of the Atkins, Ornish, Weight Watchers, and Zone diets showed similar modest reductions in weight and cardiac risk factors at 1 year. Importantly , all diets had adherence rates of only 50% to 65%, and adherence, more than macronutrient composition of the diets, had a significant correlation with weight loss. ⁸⁰

Pharmacological therapies for weight loss include sympathomimetic drugs (eg, sibutramine), drugs that alter fat digestion (eg, orlistat), antidepressants, antiepileptics, and diabetes drugs (eg, metformin). Many of these have shown modest efficacy for weight loss over placebo. A meta-analysis of sibutramine demonstrated additional mean weight loss of 4.5 kg over placebo. ⁸¹ Similarly, a meta-analysis of orlistat showed an additional mean weight loss of 2.89 kg. ⁸² As with all medications, adherence is essential for maximum efficacy of therapy, and these weight loss medications require at least 1 year if not 2 years of therapy. Tellingly, only 67% of the participants in the Kelley orlistat trial completed therapy.

Finally, an invasive option for weight loss is bariatric surgery. The procedure is generally successful; a meta-analysis -demonstrated that mean weight loss is 61% of baseline weight. ⁸³ In addition, cardiac risk factors such as hypertension and diabetes were greatly improved in the majority of patients. Despite these encouraging results, the surgery has 30-day mortalities ranging from 0.1% to 1%. In addition, the salutary -effects on weight loss and cardiac risk factor control are primarily seen only in morbidly obese patients, making this therapy less attractive for less obese patients.

FACTORS ASSOCIATED WITH NONADHERENCE

Treatment adherence is a multidimensional construct that is determined by the interplay of a variety of factors. However, many providers believe that patients are solely responsible for their adherence behavior and thus fail to account for other factors outside a patient's locus of control that can facilitate or impede the ability to adhere to a prescribed treatment (ie, therapy or system factors). ⁸⁴ Understanding of these factors and their impact on adherence behavior is essential to the proper design of interventions to improve adherence rates.

The 2003 World Health Organization report Adherence to Long-Term Therapies: Evidence for Action organized the factors affecting adherence into five dimensions: social/economic factors, health care system and team-related factors, condition-related factors, therapy-related factors, and patient-related factors (Fig. 35-2). 84 The bulk of prior research on adherence factors has focused primarily on patient-related factors, but there is increasing recognition, with accompanying research, that factors in the other four dimensions are equally important in characterizing and improving adherence.

Socioeconomic factors are major determinants of a patient's ability to adhere to treatment plans. Several studies have documented that low socioeconomic status, financial difficulty, high cost of medications, low health literacy, and unemployment are correlated with poor treatment adherence. ^{85,86} These factors are especially onerous in the treatment of chronic cardiac risk factors, for which multiple medications are required to be administered for years.

Health care system and team-related factors are another important component in treatment adherence. The currently fragmented health care system in the United States presents multiple challenges in maintaining good treatment adherence . Imposed limits on access to care, restricted medication formularies, switching of formularies, and high costs of

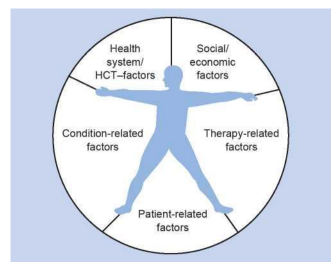


FIGURE 35-2 The five dimensions of adherence. HCT, health care team. (From Adherence to long-term therapies: evidence for action, Geneva, 2003, World Health Organization.)

medications have all been demonstrated to have a negative impact on adherence. ²

Multiple components of the individual patient-provider relationship can positively or negatively affect adherence. Bodenheimer, 87 in a recent review, described the difficulties that health care providers have in making sure their patients understand the care they receive and assisting patients in selfmanagement of their treatments. Both of these areas require attention to optimize adherence. In a review, Osterberg and Blaschke ² outlined provider actions that have a negative impact on subsequent treatment adherence, including the prescription of complex dosing regimens, the failure of the provider to explain the benefits and potential side effects of treatments, the failure of the provider to consider the patient's lifestyle or the affordability of medications in prescription decisions, and merely the presence of a poor provider-patient therapeutic relationship. A series of studies looking at patient understanding of treatment plans under scores these points. Roter and Hall 80 found that 50% of patients leave an office not understanding what they were told by the provider. Schillinger and colleagues, 89 on asking patients to restate the provider's instructions, found that 47% responded incorrectly. Finally, in another study by Schillinger and colleagues, 90 50% of patients, when asked to state how they were supposed to take a prescribed medication, did not understand how the provider had prescribed the medication. Clearly, these factors will adversely affect treatment adherence.

Condition-related factors have another component affecting treatment adherence. Cardiac risk factors, as previously noted, are often asymptomatic conditions, a characteristic that obviates some of the immediate positive feedback that symptomatic conditions provide for treatment adherence (eg, analgesics for pain). In addition, the chronic nature of the conditions, often requiring daily, life-long therapy, often results in poorer adherence patterns, as the commitment to administer daily treatments (both pharmacologic and lifestyle modification) wanes.

Therapy-related factors are a major contributor to treatment adherence. Factors such as drug tolerability, regimen complexity, frequency of dosing, number of concurrent medications , and changes in medications can affect a patient's willingness and ability to adhere to treatments. ⁸⁴ A variety of studies have demonstrated an inverse correlation between treatment adherence and number of daily doses of a

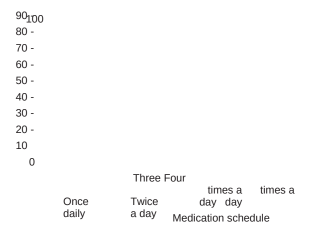


FIGURE 35-3 Adherence to medication according to frequency of doses. Vertical lines represent 1 SD on either side of the mean rate of adherence (horizontal bars). (From Osterberg L, Blaschke T: Adherence to medication. N Engl J Med 353:487, 2005.)

medication prescribed (Fig. 35-3). 91-94 Although some studies have found a positive correlation between number of medications and treatment adherence, 95 the bulk of the evidence demonstrates that simplifying medication regimens as much as possible increases the likelihood of adherence. 96

Finally, patient factors are an important aspect of treatment adherence. Interestingly, basic demographic categories - age, sex, race, and educational level-are not strong predictors of treatment adherence. 97 A study by Steiner and associates 98 underlined this point by demonstrating that prediction models for treatment adherence using only patient factors were poor. In two different health systems, the C-statistics, which measure the ability of factors to distinguish between adherents and nonadherers, were only 0.58 and 0.61. On the other hand, mental health status, including depression, cognitive impairment, anxiety, stress, and substance abuse, is strongly linked to nonadherence. 2.99-101

In contrast to basic demographic patient factors, patients' knowledge about their cardiac risk factors, health beliefs, health attitudes, self-management skills, and active participation in their care appear to be strong predictors of adherence. 100,102-104 Several studies have documented that patient beliefs and attitudes, such as low perceived risk of a condition, low perceived need for treatment, perceived ineffectiveness of a treatment, or perceived harm from a treatment, negatively affect adherence. 100,105,106 These beliefs are especially prevalent in asymptomatic conditions such as cardiac risk factors, underlining the challenge to treatment adherence for these conditions. Other patient characteristics that negatively affect adherence include a lack of self-efficacy, negative or avoidant coping strategies, and different degrees of readiness to change. 100,105,106 Patients' misperceptions or lack of insight into their particular condition or conditions can also negatively affect adherence. 107-109 For example, a common patient belief about hypertension is that it is an intermittent but acutely symptomatic condition that can be treated solely with nonpharmacologic therapies - beliefs that would seem to invalidate the need for lifelong adherence to hypertension treatments. 110-112

INTERVENTIONS TO IMPROVE ADHERENCE

Once gaps in adherence and the factors underlying them are understood, interventions can be designed to improve them. However, successful interventions to improve adherence are rare and point to the difficulty in overcoming the multiple barriers to adherence in the treatment of cardiac risk factors. Because the factors that result in nonadherence are multiple and involve patient, provider, and health care system factors, interventions that are successful and sustainable will need to be multimodal, to account for the particular barriers to adherence in their targeted population, and to integrate well into practice patterns.

Several Cochrane reviews have been conducted to assess the overall effectiveness of interventions to improve treatment adherence for chronic cardiac risk factor conditions. 113,114 In general, synthesis of common elements of effective interventions was difficult because of the multifaceted nature of many of the interventions. Nevertheless, behavioral interventions, such as reducing dosing demands or incorporating monitoring and feedback to patients and providers, had the most successful, albeit modest, effects. 115

In general, elements of successful interventions can be divided into patient, provider, and health care system factors. In a review of patient-focused interventions, the most effective strategy was simplifying the treatment regimen. 114 In contrast, motivational strategies, such as home monitoring, small group sessions, or reminder calls, were only moderately effective, and patient education alone was least effective. Claxton and 35 colleagues 8 demonstrated that simplifying medication dosing by reducing the frequency of daily medication administration was associated with improved adherence. Additional studies supported this conclusion and found that minimizing the total number of daily doses of medication was more effective at improving adherence than minimizing the total number of medications. 93,116 In addition to minimizing daily doses, selection of medications with longer half-lives, and thus "more forgiving" of occasional nonadherence, may improve overall cardiac risk factor control, such as with hypertension. 6,7 Monitoring and feedback, although less effective than medication simplification, can nevertheless be valuable in improving adherence. 117,118 For example, a trial using MEMS caps to inform adherence behavior and to provide feedback to both patients and providers among patients with refractory hypertension demonstrated improved adherence. 119 Among these patients, 30% had improvement in adherence and blood pressure control with monitoring; 20% had treatment nonadherence identified; and a subgroup of patients achieved blood pressure control with audit, feedback, and medication adjustment with use of the MEMS information.

In contrast to patient-focused interventions, provider-focused interventions have been largely disappointing. Continuing medical education efforts, audit and feedback programs, academic detailing, and computerized decision support have all been ineffective at improving adherence. 120 Interestingly, one of the few successful provider-focused interventions targeted the specific interaction between providers and patients. Rosen and colleagues 121 demonstrated improved adherence among diabetic patients by use of feed back from MEMS caps data as the basis for a conversation between providers and patients on barriers to adherence and potential solutions (eg, establishing cues to trigger timely medication administration).

Health care system interventions have had success in improving adherence when team management approaches for risk factor management were employed. 122,123 In Cooper's review of successful cardiac risk factor interventions, seven of the eight successful programs used either nurse- or pharmacist-directed programs or collaborative care of the patient with a pharmacist. ¹²⁰ In addition, employment of communication with patients between clinic visits, by modalities such as the telephone or the Internet, has also demonstrated promise. Piette and colleagues 124 showed improved adherence to diabetic treatments with a



FUTURE DIRECTIONS IN ADHERENCE

To date, most of the research into cardiac risk factor treatment nonadherence has focused on the patient, provider, health care system, condition, and treatment factors associated with nonadherence. The next step, then, is to translate these insights into effective interventions to improve adherence. Research into both the barriers to adherence and successful adherence interventions suggests that fruitful paths for improvement may involve teambased approaches to risk factor management; movement from CONCLUSION episodic, office-based management of cardiac risk factors to more changes in care.

35 locus of risk factor management from physicians and assign simple interventions have been elusive. Nevertheless, promising composed of nurses and pharmacists. This redesign overcomes the team based management, and patient activation, have been multiple barriers that physicians face in attempting to manage identified and can serve as a foundation for larger, multimodal chronic conditions in the compressed setting of a short office visit — interventions that can improve adherence and realize the ultimate competing demands, "tyranny of the urgent," and clinical inertia. In goal of optimal risk factor control and minimization of adverse addition, nurses and pharmacists who are charged with risk factor cardiac outcomes among the CAD population. management, by virtue of having a more single-minded focus to their efforts, can achieve an efficiency and experience in identifying REFERENCES nonadherence among the cardiac population and methods to overcome it.

Movement of risk factor management and adherence efforts from the office to the home setting can potentially improve adherence. Both blood pressure and lipid measurements can occur more frequently between episodic office visits, and information can be relayed to both providers and patients by telephone, mail, and Internet. In addition, information about medication refill patterns and other adherence data can also be fed back to providers and patients in a "real time" format. This iterative audit and feedback mechanism for both risk factor control and adherence can allow positive reinforcement of treatment, which is an especially important feature for chronic, asymptomatic conditions. In addition, this process can identify problems earlier and allow corrective measures to occur sooner, thus minimizing the total amount of time of nonadherence and suboptimal control of risk 10. factors.

Patient activation is another potentially important area for 11. adherence improvement. Improving a patient's understanding of the condition, the need for treatment, and the effectiveness of controlling risk factors can lead to improved "buy-in" of treatment 13. and subsequent adherence. In addition, these tar geted patient education efforts could be combined with a personalized 14. assessment of each patient's lifestyle to identify potential barriers to adherence, to direct inquiry to patients about their adherence behavior, and to identify solutions before significant nonadherence 16.

Finally, health care systems will need to provide incentives and to remove disincentives for measures to improve adherence. 18. Improved access to medical care and medications will need to 19. allow efficient management of medication and visit costs. In addition, improved delivery of medications, such as automating refills and allowing mail delivery of medications, may also facilitate 21. DiMatteo MR, Giordani PJ, Lepper HS, Croghan TW: Patient adherence and medical treatment adherence. For more continuous, home-based care to be feasible, improvements in communication between patients and health care 22. teams will be necessary. Although privacy issues will need consideration, tools such as automated telephone calls and Internet-based communication could be employed to assist with this process.

Another potential system feature that could positively affect medication adherence is automated screens of patient populations

notification of missed medication refills) and cardiac risk factor control. Such screens necessitate electronic medical records with good data quality but could potentially result in early identification of and intervention for patients with suboptimal adherence and control. Finally, any system redesign will require alignment of reimbursement to facilitate its success. Because most medical care is currently reimbursed in fee-for-service, office- or procedurebased model, changes that move care to a team-based and homebased model will need accompanying changes to ensure fair reimbursement for these efforts.

continuous, home- based management; activation of patients in Treatment adherence is an essential component to optimize cardiac their care; and alignment of health care systems to facilitate these risk factors, and significant gaps currently exist in its provision. Causes of these gaps are a complex interplay of patient, provider, The majority of successful adherence interventions remove the health care system, condition, and treatment factors. Accordingly, responsibility to a team of other health care providers, usually elements to improve adherence, such as medication simplification,

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CHAPTER 36

Clinical Practice Guidelines and Performance Measures in the Treatment of Cardiovascular Disease

Sidney C. Smith, Jr.

KEY POINTS

- Therapeutic options for treatment of patients to prevent cardiovascular disease events have increased at an exponential rate during the past quarter century.
- The Institute of Medicine has emphasized that treatment strategies and recommendations must be founded on a strong evidence base.
- Comprehensive scientific guidelines, performance measures, and policies supporting preventive therapies will rely on this foundation of scientific information.
- Evolving data confirm the efficacy of evidence-based medicine to improve patient outcomes
- It is imperative that health care providers shape their practices to keep abreast of the concepts put forth in the cardiovascular disease guidelines and performance measures.

The therapeutic options for treatment of patients to prevent cardiovascular disease (CVD) events have increased at an exponential rate during the past quarter century. This increase in medical therapies, procedures, and tests coupled with growing numbers of patients at risk for CVD events or surviving after an event has resulted in a wide array of treatment strategies, often leaving the health care provider uncertain about the appropriate course of action. Mounting evidence during the past decade has documented significant variation in the performance of recommended tests and therapies and patient outcomes. 1,2 On the basis of these trends and observations, the Institute of Medicine recommended in its 2001 report Crossing the Quality Chasm: A New Health System for the 21st Century, ³ "Patients should receive care based on the best available scientific knowledge. Care should not vary illogically from clinician to clinician or from place to place." The car diovascular responded has community by development of evidence-based practice guidelines and performance measures with the intent of providing consistent high-quality care and outcomes for patients with CVD.

CLINICAL PRACTICE GUIDELINES: BACKGROUND

Current strategies for prevention or treatment of CVD are increasingly based on guideline recommendations. Among such guideline statements, those issued by the American College of Cardiology (ACC) and the American Heart Association (AHA) have gained widespread recognition and use. ^{4,5}The origins of the ACC/AHA practice guidelines date to the early 1980s, when the federal government requested the recommendations of the ACC and AHA on the indications for cardiac pacemakers. In the early years of the guideline efforts, several statements were developed for a variety of procedures and diagnostic tests, including coronary bypass

graft surgery, exercise treadmill testing, percutaneous coronary interventions, and radionuclide testing. Understanding physicians providing patients who need the information on treatment management of a complete disease state rather than isolated advice about when to perform a procedure or diagnostic test, the ACC/AHA practice guidelines have evolved to a focus on disease states such as ST-segment elevation myocardial infarction (STEMI), valvular heart disease, chronic stable angina, and heart failure. This has resulted in an increasing number of recommendations supported by varying levels of evidence and expert consensus opinion.

GUIDELINE RECOMMENDATIONS AND EVIDENCE

The definitions used by the ACC/AHA guideline statements for the classification of recommendations and supporting level of evidence are shown in Figure 36-1. The recommendations range from Class I, for which the benefit of a test, treatment, or procedure greatly exceeds the risk and it is recommended that the test, treatment, or procedure be done, to Class III, for which the benefit of a test, treatment, or procedure is less than or equivalent to the risk and there fore should not be done. Situated between these strong recommendations in favor for or against performing a test, procedure, or treatment are Class IIa and IIb recommendations, for which benefit exceeds risk such that the performance of the test, procedure, or treatment is deemed reasonable (IIa) or might be considered (IIb). For each class of recommendation, the guidelines must further state the level of evidence on which the recommendation is based. Level A represents multiple randomized clinical trials or metaanalyses obtained in multiple subgroups including age, gender, and ethnicity;



Benefit > Risk Benefit >>>Risk Benefit >>Risk Risk > Benefit Additional studies with focused Additional studies with broad Procedure/Treatment Procedure/Treatment should SHOULD be objectives needed; additional objectives needed performed/administered IT IS REASONABLE to registry data would be helpful performed/administered SINCE perform procedure/administer Procedure/Treatment IT IS NOT HELPFUL AND MAY treatment **MAY BE CONSIDERED** BE HARMFUL LEVEL Multiple populations Recommendation that Recommendation in favor of Recommendation that procedure or treatment is evaluated* treatment or procedure being Recommendation's procedure or treatment is not Data derived from multiple useful/effective useful/effective usefulness/efficacy less well useful/effective and may be randomized clinical trials or established harmful ■ Sufficient evidence from ■ Some conflicting evidence meta-analyses multiple randomized trials or from multiple randomized trials Greater conflicting evidence Sufficient evidence from meta-analyses or meta-analyses from multiple randomized trials multiple randomized trials or or meta-analyses meta-analyses Recommendation that Recommendation that ■ Recommendation in favor of ■ Recommendation's procedure or treatment is not LEVEL B procedure or treatment is treatment or procedure being usefulness/efficacy less well useful/effective and may be Limited populations evaluated* useful/effective useful/effective established Data derived from a single ■ Evidence from single Some conflicting evidence ■ Greater conflicting evidence randomized trial or ■ Evidence from single randomized trials or from single randomized trials from single randomized trials nonrandomized studies randomized trials or nonrandomized studies or nonrandomized studies or nonrandomized studies nonrandomized studies Recommendation that ■ Recommendation in favor of ■ Recommendation's Recommendation that LEVELC procedure or treatment is treatment or procedure being usefulness/efficacy less well procedure or treatment is not Very limited populations useful/effective useful/effective established useful/effective and may be evaluated* Only expert opinion, case Only consensus opinion of Only diverging expert Only diverging expert studies, or standard of care experts, case studies, or opinion, case studies, or opinion, case studies, or Only expert opinion, case standard of care standard of care standard of care studies, or standard of care Suggested phrases for shouldnt is reasonable may/might be considered is not recommended writing recommendations is recommended can be useful/effective/beneficial may/might be reasonable is not indicated should not is indicated is probably recommended usefulness/effectiveness is is not useful/effective/beneficial is useful/effective/beneficial or indicated unknown/unclear/uncertain may be harmful or not well established

CLASS IIa

FIGURE 36-1 ACC/AHA guideline classification of recommendations and level of evidence

Data available from clinical trials or registries about the usefulness/efficacy in different subpopulations, such as gender, age, history of diabetes, history of prior myocardial infarction, history of heart failure, and prior aspirin use. A recommendation with Level of Evidence B or C does not imply that the recommendation is weak. Many important clinical questions addressed in the guidelines do not lend themselves to clinical trials. Even though randomized trials are not available, there may be a very clear clinical consensus that a particular test or therapy is useful or effective.

CLASS IIb

CLASS III

* In 2003, the ACC/AHA Task Force on Practice Guidelines developed a list of suggested phrases to use when writing recommendations. All recommendations in this guideline have been written in full sentences that express a complete thought, such that a recommendation, even if separated and presented separately from the rest of the document (including headings above sets of recommendations), would still convey the full intent of the recommendation. It is hoped that this will increase readers' understanding of the guidelines and will allow queries at the individual recommendation level. (From Smith SC, Allen J, Blair SN, et al: AHA/ACC guidelines for secondary prevention for patients with coronary and other atherosclerotic vascular disease: 2006 update: endorsed by the National Heart, Lung, and Blood Institute. Circulation 113:2363, 2006.)

level B represents data from single randomized trials or non- - relationships are present, and their recusal is noted in the guideline randomized studies in which limited populations have been statement. Membership of the guideline writing groups is being evaluated; and level C is based on consensus opinion of experts, structured with a goal that no more than 30% of the writing group case studies, or standards of care. A level of evidence of B or C does may have conflicts of interest or relationships with industry for the not necessarily imply a weak recommendation. There are many tests, procedures, and treatments being discussed. Further review important clinical questions for which the number of patients and recommendations for approval of all ACC/AHA guidelines available for a randomized controlled trial is too small to provide are provided by additional individuals, including members of the significant results or the clinical circumstances are such that there Science Advisory Committee of the AHA, Board of Trustees of the is a clear and strong consensus regarding the indication for a test, ACC, expert content reviewers, and representatives from other procedure, or treatment. However, the guidelines are written with societies participating in the the objective of having the strongest possible evidence base.

CLASS I

During recent years, considerable efforts have been directed towards avoiding conflicts of interest and relation ships with industry that might potentially bias the recommendations in guideline statements. The ACC and AHA have instituted strong policies to see that such interests are avoided in the development of guideline statements. Members of guideline writing committees must recuse themselves from voting when such conflicts or

development or endorsement of the guideline. These measures recommendation is of lower certainty, such as Class IIb, and 581 help ensure that the guidelines are strongly based on evidence or objective expert consensus.

STATUS OF CURRENT GUIDELINES **EVIDENCE BASE**

The largest continuous experience with the development of cardiovascular guidelines is found in the joint efforts of the ACC and AHA. 4.5 These guidelines have provided critical information to assist with the establishment of standards of care and benchmarks to assess the quality of care. The evolution of recommendations in these guidelines and the development of evidence base for these recommendations have been analyzed and provide insight into challenges facing development of current guideline statements. 6 Since the first ACC/AHA clinical practice guideline was released in 1984 on the indications for cardiac pacemakers to September 2008, the ACC/AHA Task Force on practice guidelines has published 53 guidelines on 22 topics resulting in a total of 7196 recommendations. Among these 53 guidelines, there were 22 that were disease based, 15 that were interventional procedure based, and 14 that were diagnostic procedure based. The use of level of evidence for guideline recommendations was first introduced in 1998, and analysis of the 16 guidelines current at the time of the report 6 comprising a total of 2711 recommendations revealed a high proportion of recommendations (48%) to be based on expert consensus, standard of care, or case studies (level of evidence C), whereas a small number (11%) were derived from a high evidence base of multiple randomized controlled trials or metaanalyses (level of evidence A). Moreover, among those guidelines that had undergone one revision or update, there was a 48% increase in recommendations, with the greatest number occurring in Class II, indicating a level of uncertainty in the recommendation . Importantly, the secondary prevention guideline recommendations had the highest evidence base, with more than 90% being level of evidence A or B, attesting to the strong evidence available to support recommendations for preventive therapies.

FUTURE CONSIDERATIONS FOR GUIDELINE DEVELOPMENT

The observations from the review of ACC/AHA guidelines support the need for more evidence to assist with the clinical decisions made by physicians who treat patients with CVD. The current system of clinical research that generates evidence from randomized controlled trials is largely supported by the pharmaceutical industry and in most instances understandably directed towards gaining approval of new medical therapies. There is limited sponsorship of trials that focus on questions relating to clinical practice combining or comparing existing tests, procedures, and treatments. Broader support and funding are necessary if we are to advance the knowledge base for guideline development.

The purpose of the ACC/AHA practice guidelines is to assist health care providers in clinical decision making by describing a range of reasonably acceptable approaches to assist in the diagnosis, management, and prevention of cardiovascular diseases or conditions. Guideline recommendations attempt to define practices that will meet the needs of most patients in most circumstances. However, the ultimate judgment regarding the care of a particular patient should be made by the health care provider and patient, with all of the circumstances presented by that patient taken into consideration . Thus, there are circumstances in which deviations from these guidelines may be appropriate, especially when a

supported primarily by expert opinion level C. Clinical decision making should also consider the quality and availability of expertise in the area where care is provided. On occasion, the ACC/AHA guidelines may be used as the basis for regulatory or payer decisions, but their ultimate goal is to improve the quality of care and to serve the patient's best interests.

GUIDELINES FOR SECONDARY PREVENTION

The AHA/ACC Guidelines for Secondary Prevention, endorsed by the National Heart, Lung, and Blood Institute (NHLBI), provide concise evidence-based recommendations for the prevention of cardiovascular events in patients with established CVD. 7 The goal for controlling each risk factor is listed in conjunction with recommended therapeutic interventions to achieve that goal (Table 36-1). The class of recommendation and evidence base for each intervention are concisely summarized, and the references and supplementary search criteria are provided. As noted earlier in this chapter, these guidelines have a high evidence base, with more than 90% being level A or B. They are widely used and quoted and provide the basis for the They are widely used and quoted and provide the basis for the **S** AHA and ACC quality improvement programs reviewed later **S** in this chapter.

The recommendations are closely coordinated with statements from the National Institutes of Health/NHLBI and the Centers for Disease Control and Prevention (CDC) and the Centers for Disease Control and Prevention (CDC) and updated on the basis of the evolving evidence base for each risk factor. For example, recommendations on target goals for lipids and hypertension are consistent with those put forth by the NHLBI National Cholesterol Adult Treatment Panel 8,9 and the Joint National Committee on Prevention, Detection, Evaluation, and Treatment of High Blood Pressure, 10 and the recommendations for influenza vaccine in all patients with atherosclerotic vascular disease are consistent with those of the CDC Advisory Committee on Immunization Practices. ¹¹ These guideline recommendations are also integrated into other 8 relevant guideline statements from the ACC/AHA, such as those for ST-segment myocardial infarction, unstable angina/non-ST-segment myocardial infarction, chronic stable angina and percutaneous coronary interventions, and undated angina, and percutaneous coronary interventions, and updated as dictated by new evidence from randomized controlled trials and other sources.

A consistent focus of these guidelines has been the assessment of the evidence base according to gender, age, and ethnicity. Many randomized controlled trials have cut points for study inclusion at 75 or 80 years of age or have under representation of women or minority ethnic groups. Thus, the evidence base for recommendations may not be universally consistent across age, gender, and ethnic groups. Recognizing the differences that do exist in the evidence base for women and the elderly, the AHA has developed guideline statements 12,13 focusing specifically on the evidence base for those groups that serve as valuable additions to the general guide line statements.

PERFORMANCE BACKGROUND

MEASUREMENTS:

Numerous studies have demonstrated a gap between guideline recommendations and the quality of care rendered to patients. The Institute of Medicine has outlined deficiencies that exist in the delivery of effective, timely, safe, and equitable care to patients. Recognizing the importance of these concerns, the ACC and AHA have taken a leadership role in the development of performance measures 14 for several cardiovascular conditions (Table 36-2). These performances

Goal

Complete cessation. No exposure to environmental tobacco smoke.

Blood Pressure Control

Goal

<140/90 mm Hg

<130/80 mm Hg if patient has diabetes or chronic kidney disease

Lipid Management

Goal

If triglycerides are >200 mg/dL, non-HDL-C should be <130 mg/dL-

Intervention Recommendations with Class of Recommendation and Level of Evidence

- Ask about tobacco use status at every visit. I (B)
- Advise every tobacco user to guit. I (B)
- Assess the tobacco user's willingness to quit. I (B)
- Assist by counseling and developing a plan for guitting. I (B)
- Arrange follow-up, referral to special programs, or pharmacotherapy (including nicotine replacement and bupropion). I

(B)

Urgent avoidance of exposure to environmental tobacco smoke at work and home. I (B)

For all patients:

· Initiate or maintain lifestyle modification - weight control; increased physical activity; alcohol moderation; sodium reduction; and emphasis on increased consumption of fresh fruits, vegetables, and low-fat dairy products. I (B)

For patients with blood pressure > 140/90 mm Hg (or > 130/80 mm Hg for individuals with chronic kidney disease or diabetes): · As tolerated, add blood pressure medication, treating initially with beta blockers and/or ACE inhibitors, with the addition of other drugs such as thiazides as needed to achieve goal blood pressure . I (A)

[For compelling indications for individual drug classes in specific vascular diseases, see Seventh Report of the Joint National Committee on Prevention, Detection, Evaluation, and Treatment of High Blood Pressure (JNC 7).]

For all patients:

- Start dietary therapy. Reduce intake of saturated fats (to < 7% of total calories), trans-fatty acids, and cholesterol (to < 200 mg/d). I (B)
 - Adding plant stanol/sterols (2 g/d) and viscous fiber (> 10 g/d) will further lower LDL-C.
 - Promote daily physical activity and weight management. I (B)
- Encourage increased consumption of omega-3 fatty acids in the form of fish or in capsule form (1 g/d) for risk reduction. For the treatment of elevated triglycerides, higher doses are usually necessary for risk reduction. IIb (B)

Assess fasting lipid profile in all patients, and within 24 hours of hospitalization for those with an acute cardiovascular or coronary event. For hospitalized patients, initiate lipid-lowering medication as recommended below before discharge according to the following schedule:

- LDL-C should be < 100 mg/dL I (A), and
- Further reduction of LDL-C to < 70 mg/dL is reasonable. IIa (A)
- If baseline LDL-C is > 100 mg/dL, initiate LDL-lowering drug therapy. \S I (A)
- If on-treatment LDL-C is > 100 mg/dL, intensify LDL-lowering drug therapy (may require LDL-lowering drug combination 11). I (A)
 - If baseline LDL-C is 70 to 100 mg/dL, it is reasonable to treat LDL-C < 70 mg/dL. IIa (B)
 - If triglycerides are 200 to 499 mg/dL, non-HDL-C should be < 130 mg/dL. I (B), and
 - Further reduction of non-HDL-C to < 100 mg/dL is reasonable. IIa (B)
 - Therapeutic options to reduce non-HDL-C are:
 - More intense LDL-C-lowering therapy I (B), or
 - Niacin ^ (after LDL-C-lowering therapy) IIa (B), or
 - Fibrate therapy # (after LDL-C-lowering therapy) IIa (B)
- If triglycerides are > 500 mg/dL, # therapeutic options to prevent pancreatitis are fibrate ^or niacin ^before LDL-lowering therapy; and treat LDL-C to goal after triglyceride-lowering therapy. Achieve non-HDL-C < 130 mg/dL if possible. I (C)
- For all patients, assess risk with a physical activity history and/or an exercise test, to guide prescription. I (B)
- For all patients, encourage 30 to 60 minutes of moderate-intensity aerobic activity, such as brisk walking, on most, preferably all, days of the week, supplemented by an increase in daily lifestyle activities (eg, walking breaks at work, gardening, household work). I (B)
 - Encourage resistance training 2 days per week. IIb (C)
- Advise medically supervised programs for high-risk patients (eg, recent acute coronary syndrome or revascularization, heart failure). I (B)

LDL-C <100 mg/dL

Physical Activity

Goal

30 minutes, 7 days per week (minimum 5 days per week)

Weight Management

Body mass index: 18.5 to 24.9 kg/m² Waist circumference: men <40 inches, women <35 inches





Assess body mass index and/or waist circumference on each visit and consistently encourage weight maintenance/reduction through an appropriate balance of physical activity, caloric intake, and formal behavioral programs when indicated to maintain/achieve a body mass index between 18.5 and 24.9 kg/m². I (B)

If waist circumference (measured horizontally at the iliac crest) is > 35 inches in women and > 40 inches in men, initiate lifestyle changes and consider treatment strategies for metabolic syndrome as indicated. I (B)

The initial goal of weight loss therapy should be to reduce body weight by approximately 10% from baseline. With success, further weight loss can be attempted if indicated through further assessment. I (B)

TABLE 36-1 AHA/ACC Secondary Prevention for Patients with Coronary and Other Vascular Disease * : 2006 Update—cont'd

Intervention Recommendations with Class of Recommendation and Level of Evidence

Diabetes Management

Goal

HbA1c < 7%

Antiplatelet Agents/Anticoagulants

- · Initiate lifestyle and pharmacotherapy to achieve near-normal HbA1c. I (B)
- B ^e g ⁱⁿ vigorous modification of other risk factors (eg, physical activity, weight management, blood pressure control, and cholesterol management as recommended above). I (B)
- · Coordinate diabetic care with patient's primary care physician or endocrinologist. I (C)
- · Start aspirin 75 to 162 mg/d and continue indefinitely in all patients unless contraindicated. I (A)
 - For patients undergoing coronary artery bypass grafting, aspirin should be started within 48 hours after surgery to reduce saphenous vein graft closure. Dosing regimens ranging from 100 to 325 mg/d appear to be effective. Doses higher than 162 mg/d can be continued for up to 1 year. I (B)
- Start and continue clopidogrel 75 mg/d in combination with aspirin for up to 12 months in patients after acute coronary syndrome or percutaneous coronary intervention with stent placement (> 1 month for bare metal stent, > 3 months for sirolimus-eluting stent, and > 6 months for paclitaxel-eluting stent). I (B)
 - Patients who have undergone percutaneous coronary intervention with stent placement should initially receive higher-dose aspirin at 325 mg/d for 1 month for bare metal stent, 3 months for sirolimus-eluting stent, and 6 months for paclitaxel-eluting stent. I (B)
- Manage warfarin to international normalized ratio = 2.0 to 3.0 for paroxysmal or chronic atrial fibrillation or flutter, and in
 post-myocardial infarction patients when clinically indicated (eg, atrial fibrillation, left ventricular thrombus). I (A)
- Use of warfarin in conjunction with aspirin and/or clopidogrel is associated with increased risk of bleeding and should be
 monitored closely. I (B)

Renin-Angiotensin-Aldosterone System Blockers

ACE inhibitors:

- Start and continue indefinitely in all patients with left ventricular ejection fraction < 40% and in those with hypertension, diabetes, or chronic kidney disease, unless contraindicated. I (A)
- · Consider for all other patients. I (B)
- Among lower-risk patients with normal left ventricular ejection fraction in whom cardiovascular risk factors are well
 controlled and revascularization has been performed, use of ACE inhibitors may be considered optional. IIa (B)

Angiotensin receptor blockers:

- Use in patients who are intolerant of ACE inhibitors and have heart failure or have had a myocardial infarction with left ventricular ejection fraction < 40%. I (A)
- · Consider other patients who have ACE inhibitor intolerance. I (B)
- · Consider use in combination with ACE inhibitors in systolic-dysfunction heart failure. IIb (B)

Aldosterone blockade:

Use in post-myocardial infarction patients, without significant renal dysfunction or hyperkalemia, who are already receiving therapeutic doses of an ACE inhibitor and beta blocker, have a left ventricular ejection fraction < 40%, and have either diabetes or heart failure. I (A)

Beta Blockers

- · Start and continue indefinitely in all patients who have had myocardial infarction, acute coronary syndrome, or left ventricular dysfunction with or without heart failure symptoms, unless contraindicated. I (A)
- Consider chronic therapy for all other patients with coronary or other vascular disease or diabetes unless contraindicated. **IIa** (C)

Influenza Vaccination

Patients with cardiovascular disease should have an influenza vaccination. I (B)

- Patients covered by these guidelines include those with established coronary and other atherosclerotic vascular disease, including peripheral arterial disease, atherosclerotic aortic disease, and carotid artery disease. Treatment of patients whose only manifestation of cardiovascular risk is diabetes will be the topic of a separate AHA scientific statement. ACE indicates angiotensin-converting enzyme.
- ^Non-HDL-C, total cholesterol minus HDL-C.
- ^Pregnant and lactating women should limit their intake of fish to minimize exposure to methylmercury.
- § When LDL-lowering medications are used, obtain at least a 30% to 40% reduction in LDL-C levels. If LDL-C <70 mg/dL is the chosen target, consider drug titration to achieve this level to minimize side effects and cost. When LDL-C <70 mg/dL is not achievable because of high baseline LDL-C levels, it is generally possible to achieve reductions of >50% in LDL-C levels by either statins or LDL-C-lowering drug combinations.
- ∥Standard dose of statin with ezetimibe, bile acid sequestrant, or niacin.
- ^The combination of high-dose statin + fibrate can increase the risk for severe myopathy. Statin doses should be kept relatively low with this combination. Dietary supplement niacin must not be used as a substitute for prescription niacin.
- # Patients with very high triglycerides should not consume alcohol. The use of bile acid sequestrant is relatively contraindicated when triglycerides are >200 mg/dL.
- Creatinine should be <2.5 mg/dL in men and <2.0 mg/dL in women.

^Potassium should be <5.0 mEq/L.

From Smith SC, Allen J, Blair SN, et al: AHA/ACC guidelines for secondary prevention for patients with coronary and other atherosclerotic vascular disease: 2006 update: endorsed by the National Heart, Lung, and Blood Institute. *Circulation* 113:2363, 2006.





Planned publication date.

SVS, Society for Vascular Surgery.

AACVPR, American Association of Cardiovascular and Pulmonary Rehabilitation; ACC, American College of Cardiology; ACCF, American College of Cardiology Foundation; ACR, American College of Radiology; AHA, American Heart Association; PCPI, American Medical Association-Physician Consortium for Performance Improvement; SCAI, Society for Cardiac Angiography and Interventions; SIR, Society for Interventional Radiology; SVM, Society for Vascular Medicine; SVN, Society for Vascular Nursing;

From Redberg RF, Benjamin E, Bittner V, et al: ACCF/AHA 2009 performance measures for primary prevention of cardiovascular disease in adults. *J Am Coll Cardiol* 54:1364, 2009.

measures can be used for quality improvement programs directed towards improving patient outcomes. The basis for performance measures has generally been Class I or Class III recommendations from the ACC/AHA guidelines, which identify procedures or treatments that should or should not be given to patients. These processes of care measures are now combined with recommendations regarding the structure of care that deal with the environment in which patients are treated and outcomes of care that assess health care systems on the basis of improved outcomes for patient care. The initial purpose of the ACC/AHA performance measures was for use in quality improvement efforts; however, their application has been expanded by other organizations for public reporting or external review. This development has led the performance measure writing groups to characterize their recommendations as either performance measures or test measures. The distinction between two is that performance measures are those recommendations that are deemed appropriate for use in both quality improvement programs and external reporting, whereas test measures are those thought to be appropriate for quality improvement but not for external reporting until their efficacy has been validated. Thus, field testing of performance measures is strongly recommended if they are considered for use beyond quality improvement.

DEVELOPMENT AND SELECTION OF PERFORMANCE MEASURES

Performance measures generally target a specific patient population that is observed during a particular time period. For example, the performance measures for primary prevention outlined later in this chapter define the population as those patients older than 18 years to avoid conflict with recommendations for children. In addition, certain measures are given an upper age limit of 80 years because of the absence of evidence to support the recommendation in patients older than 80 years. Criteria for final selection of performance measures include (1) those Class I or III guideline recommendations with the strongest class of recommendation and level of evidence, (2) the ease or complexity of the measurement, and (3) coverage in

other performance measurement sets. The phrasing of performance measures is given careful consideration . For example, many patients are unable to achieve recommended blood pressure control because of medication noncompliance, costs, side effects, or other reasons. To avoid penalizing the physician for these issues that are beyond the physician's control, the measure is written to accept documentation of the use of at least two medications in patients whose blood pressures are greater than the target of 140/90 mm Hg or the achievement of the recommended target outcome.

Potential performance measures are developed with the use of criteria that carefully define measurement parameters and evidence for or against the recommended measure as well as their applicability, interpretability, and feasibility. Specific attributes of highly desirable performance measures have been carefully defined by the ACC/AHA Task Force on Performance Measures (Table 36-3). Not all performance measures are based on level A evidence. For example, in dealing with a recommendation to counsel regarding cessation of tobacco use, the absence of multiple large randomized controlled studies does not preclude consideration of this important measure. By contrast, in the case of pharmaceutical agents that lend themselves to large randomized controlled trials, the need for level A evidence base is greater.

PERFORMANCE MEASURES FOR PRIMARY PREVENTION OF CARDIOVASCULAR DISEASES IN ADULTS

The 2009 ACC/AHA performance measures for primary prevention in adults ¹⁵ list 13 measures including recommendations involving lifestyle, blood pressure, lipids, global risk, and aspirin (Table 36-4). These 13 measures apply to all adults without CVD, including those without diabetes. They support practices expected to reduce the long-term risk of cardiovascular events. Because any single visit may not provide full opportunity for the full range of the performance

Attribute Evidence based
Interpretable Actionable
Denominator precisely defined Numerator precisely defined Validity type Face
· Content-
• Construct•
Reliability
Feasibility Reasonable effort Reasonable cost Reasonable time period for collection
Overall assessment of measure for inclusion in measurement set

[•]The measure intuitively appears to capture what it is intended to capture.

From Redberg RF, Benjamin E, Bittner V, et al: ACCF/AHA 2009 performance measures for primary prevention of cardiovascular disease in adults. *J Am Coll Cardiol* 54:1364, 2009.

measures to be addressed, at least two encounters during a period of 1 year are recommended before the physician is expected to have responsibility for primary CVD prevention. Two of the measures, aspirin use and global risk estimation, are considered appropriate for internal quality improvement only. Statins are recognized as the mainstay of lipid-lowering therapy; however, given the variable response to lipid-lowering agents and their side effects, the writing group chose to accept either the attainment of a target goal for lipid lowering or the documented use of more than one lipid-lowering agent.

VALIDATION THAT GUIDELINES AND PERFORMANCE MEASURES IMPROVE CARDIOVASCULAR DISEASE OUTCOMES

The AHA and ACC have both developed programs aimed towards improving the use of guideline-recommended therapies by physicians. The ACC Guidelines Applied in Practice (GAP) program studied the potential benefits of guideline-recommended evidenced-based therapies among 2857 Medi care patients admitted to 38 hospitals in southeastern Michigan. ¹⁶ For GAP patients receiving guideline-recommended therapies, a significant reduction was observed in the in-hospital, 30-day, and 1-year mortality. After multivariate analysis, GAP accounted for a 21% to 26% reduction in mortality.

The AHA Get With The Guidelines (GWTG) program uses local physician champions, multidisciplinary teams, and preprinted orders along with a web-based Patient Management Tool to improve the use of evidence-based guideline-recommended therapies. Early results from GWTG ¹⁷ from 45,988 patients in 92 hospitals revealed improvement in the use of 10 of 11 guideline-recommended measures. This

	ovascular Disease Performance prement Set	
Performance Measure Name	Measure Description	Designation
Lifestyle/risk factor screening	Assessment of lifestyles and risk factors for development of CVD	A/PR IQI
2. Dietary intake counseling	Counseling to eat a healthy diet	A/PR
Physical activity counseling	Counseling to engage in regular physical activity	A/PR
4. Smoking/tobacco use	Risk assessment for smoking and tobacco use behaviors	A/PR IQI
5. Smoking/tobacco cessation	Cessation intervention for active smoking (tobacco use)	A/PR
6. Weight/adiposity assessment	Measurement of weight and body mass index and/or waist circumference	A/PR
7. Weight management	Counseling to achieve and maintain ideal body weight	A/PR IQI
Blood pressure measurement	Measurement of blood pressure in all patients	A/PR
9. Blood pressure control		A/PR IQI
	Effective blood pressure control or combination therapy for patients with hypertension	
10. Blood lipid measurement	Fasting lipid profile performed	A/PR IQI
11. Blood lipid therapy and control	Proportion of patients who meet current LDL-C treatment targets or who are prescribed >1 lipid-lowering medications at maximum tolerated dose	A/PR
12. Global risk estimation	Use of a multivariable risk score to estimate a patient's absolute risk for development of coronary heart disease	IQI
13. Aspirin use	Aspirin in patients without clinical evidence of atherosclerotic disease who are at higher CVD risk	IQI

TABLE 36—4 ACC/AHA Primary Prevention of

A/PR, accountability/public reporting measures (appropriate for all uses, including internal quality improvement, pay for performance, physician ranking, and public reporting); CVD, cardiovascular disease; IQI, internal quality improvement measures (recommended for use in internal quality improvement programs only; not appropriate for any other use, eg, pay for performance, physician ranking, or public reporting); LDL-C, low-density lipoprotein cholesterol.

From Redberg RF, Benjamin E, Bittner V, et al: ACCF/AHA 2009 performance measures for primary prevention of cardiovascular disease in adults. *J Am Coll Cardiol* 54:1364, 2009

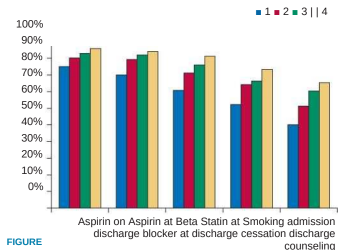
improvement in the use of recommended therapies occurred during a 1-year period, much faster than would have been expected from traditional quality improvement programs . Subsequent studies ¹⁸ comparing the use of guideline-recommended measures in 233 GWTG hospitals with 3407 non-GWTG hospitals also observed better results for the GWTG hospitals. Results from the GWTG data base have also provided important information about differences in therapy.



¹The extent to which the items comprehensively capture the domain they are intended to measure.

[•]The extent to which the measures correlate with other methods of quantifying the underlying construct.





36-2
Compliance by quartile with guidelines of acute myocardial infarction. Bars show percentage compliance for quartiles of hospitals grouped by performance. (From Braunwald's heart disease, ed 8; after Peterson ED, Roe MT, Mulgund J, et al: Association between hospital process performance and outcomes

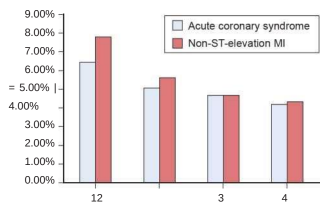
Braunwald's heart disease, ed 8; after Peterson ED, Roe MT, Mulgund J, et al: Association between hospital process performance and outcomes among patients with acute coronary syndromes. JAMA 295:1912, 2006.)

In 78,254 patients with acute myocardial infarction admitted to 420 GWTG hospitals, women were shown to be less likely to receive aspirin and beta blockers on admission and less likely to undergo cardiac catheterization and revascularization. ¹⁹

Important observational data indicate that broader use of guideline-based therapies is associated with improved outcomes. 20 Analysis of hospital care in 350 academic and non academic hospitals involving 64,775 patients presenting with findings consistent with non-ST elevation myocardial infarction revealed wide variation in adherence (63% to 82%) to nine ACC/AHA guideline-recommended therapies (Fig. 36-2). Furthermore, the guideline adherence rate was significantly associated with in-hospital mortality, with mortality rates significantly decreasing from 6.31% for the lowest adherence quartile to 4.15% for the highest adherence quartile. After risk adjustment, every 10% increase in composite adherence to guidelines at a hospital was associated with a 10% decrease in patients' likelihood of in-hospital mortality (Fig. 36-3). These findings strongly support the use of guideline-based treatments at the system-wide level for the management of patients with CVD.

ADHERENCE TO PREVENTIVES THERAPIES

The success of evidence-based guideline implementation is founded on adherence by patients and physicians to the recommended therapies. As aptly put by Surgeon General C. Everett Koop, "Drugs don't work in patients who don't take them." Effective adherence to recommended medical therapies involves an active partnership between patients and physicians within a health care system to implement and to continue appropriate treatments and behaviors. Unfortunately, there is mounting evidence that nonadherence is widely present and results in worsened outcomes and increased cost of care. 21 Nearly one of four patients with myocardial infarction does not fill medications after discharge from the hospital by day 7. 22 Lack of adherence to medications has been shown to result in adverse outcomes. 23 Among patients with chronic coronary heart disease, non- adherence to medications such as beta blockers, statins, and angiotensin-converting enzyme inhibitors has been



Quartile for hospital composite compliance with guidelines (lowest to highest)

FIGURE 36-3 In-hospital mortality for patients with acute coronary syndrome and non-ST elevation myocardial infarction correlated with hospital compliance with guidelines. Bars show percentage compliance for quartiles of hospitals grouped by performance. (From Braunwald's heart disease, ed 8; after Peterson ED, Roe MT, Mulgund J, et al: Association between hospital process performance and outcomes among patients with acute coronary syndromes. JAMA 295:1912, 2006.)

TABLE 36-5	Reasons for Medical Nonadherence				
Categories of Nonadherence Examples					
Health system					
		Poor quality of provider-patient relationship; poor communication; lack of access to health care; lack of continuity of care			
Condition Asymptomatic chronic disease (lack of physical cues); mental health disorde (eg, depression)					
Patient					
		Physical impairments (eg, vision problems or impaired dexterity); cognitive impairment; psychological/behavioral; younger age; nonwhite race			
Therapy		Complexity of regimen; side effects			
Socioeconomic		Low literacy; higher medication costs; poor social support			

From Ho PM, Bryson CL, Rumsfeld JS: Medication adherence. *Circulation* 119:3028,

associated with a 10% to 40% increase in hospitalizations for cardiovascular causes and a 50% to 80% relative increase in risk of mortality. $^{24}\,$

There are many reasons for nonadherence that have been summarized by the World Health Organization into five major categories (Table 36-5). ²⁵ Younger patients, especially those who are depressed or nonwhite, are less likely to adhere to recommended therapies. Chronic conditions requiring ongoing therapy are associated with nonadherence. Low literacy, higher medication costs, and poor social support are important considerations. Successful implementation of medical therapies requires a combined effort involving health care providers, patients, and organizations delivering or paying for health care. The AHA GWTG program has demonstrated success in improving health care provider adherence to recommended guideline therapies. Patient education is the first step to improve patient adherence and must be

repeated on subsequent outpatient visits. Telephone follow-up and other reminders can also be an effective tool to improve adherence. Finally, hospitals and payers are increasingly recognizing that the cost of medical care is adversely associated with nonadherence and are therefore becoming involved in programs that assist health care providers and patients to implement and to sustain recommended therapies.

Whereas great progress has been recognized in the development of guidelines and performance measures for patient care, the success of these efforts resides with effective programs directed towards ensuring adherence by providers and patients with recommended therapies. This will be a growing area of research and emphasis for our health care system.

CONCLUSION

Health care is currently undergoing major changes. The Institute of Medicine has emphasized that treatment strategies and recommendations must be based on a strong evidence base. Comprehensive policies supporting preventive thera pies will rely on this foundation of scientific information. Evolving data confirm the efficacy of evidence-based medicine to improve patient outcomes. It is imperative that health care providers shape their practices to keep abreast of the concepts put forth in the CVD guidelines and performance measures and enable patients to adhere to the recommended therapies.

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This page intentionally left blank studies indicating a lower risk for development of coronary heart disease (CHD) for moderate consumers of alcohol. ¹⁻⁴ Of Hun dreds of studies published in the scientific

literature during the past few decades, the results have been extremely consistent. Almost uniformly, they have demonstrated that in comparison with nondrinkers, moderate drinkers are less likely to develop CHD.

In a meta-analysis by Corrao and associates

ates ⁵ based only on high-quality prospective studies, there was a J-shaped curve for the association between alcohol and CHD, with its nadir at about 20 g/day of alcohol, the equivalent of about 1% US drinks (with

Protein kinase C serir phosphorylates insulin receptors, interfering with insulin

growth explosive The cardiovascular imaging during the past few decades has facilitated the noninvasive detection of CAD. Although the CAC score correlates well with disease burden, calcified plagues represent only a portion of the total atherosclerosis plaque burden. 15 With the advent of multiple CT technology, coronary computed tomographic angiography (CTA) has the potential provide comprehensive information about the location, severity, and characteristics of atherosclerotic plaque. It is able to differentiate plaques that are calcified, noncalcified, and mixed (containing both

American The Association recommendations for calcium scanning from 2006 Scientific Statement summarized as follows: (1) CAC scanning may be suitable in patients at intermediate CAD risk to refine risk prediction and to select patients for altered targets of lipidlowering therapies; (2) CAC assessment may be reasonable for the assessment of patients, symptomatic especially in the setting of

equivocal treadmill or functional testing; and (3) CAC scan may be considered for triage of patients with chest pain with equivocal or normal electrocardiograms and negative cardiac enzyme studies. ²⁴

Other scientific statements have also endorsed the conceptual approach to refining the cardiovascular assessment through CAC detection. For example, the NCEP ATP III stated that "in persons with multiple risk factors, high coronary calcium scores (eg, > 75th percentile for age and sex) denote advanced coronary atherosclerosis and provide a rationale for intensified LDL-lowering therapy." 8

A clinical expert consensus document of the American College of Cardiology published in 2007 specified that calcium coronary measurement in clinically selected patients at intermediate CAD risk (eg, those with a 10% to 20% Framingham 10-year risk score) is a reasonable option to refine clinical risk prediction and to select patients for altered targets for lipid-lowering therapies. recommendations from the American Heart Association and the American College of Cardiology are similar to those of the European guidelines. The European guidelines state, "The resulting calcium score is an important parameter to detect asymptomatic individuals at high risk for future CVD events, independent of traditional risk factors." The guidelines state that calcium scanning should be used as a tool to improve risk assessment in individual patients. This organization further acknowledged that the prognostic relevance of CAC has been demonstrated in several prospective studies, not only in asymptomatic individuals but also in patients undergoing coronary angiography. However, screening for CAC should be reserved for individuals at intermediate risk and in men older than 45 years and women older than 55 years.